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(54) Title: SPLICE VARIANT OF HUMAN SODIUM III CHANNEL (HNAIII18)

(57) Abstract: Described herein is a splice variant of the human NaIII channel α subunit, designated hNaIII18. Also described are nucleotide and amino acid sequence for hNaIII18, oligonucleotide primers and probes for hNaIII18, hNaIII18 regulatory sequences, hNaIII18-specific antibodies, methods of detecting hNaIII18 proteins or nucleic acids, and methods of screening for modulators of hNaIII18 expression or activity.



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Splice Variant of Human Sodium III Channel (hNaIII18)

This application claims priority from U.S. Provisional Application
Serial No. 60/431,794, filed December 4, 2002, which is hereby incorporated by
5 reference in its entirety.

FIELD OF THE INVENTION

The present invention relates to a human splice variant of the voltage-
gated sodium III channel, termed hNaIII18, as well as methods for stable expression
10 of hNaIII18 in cell lines, and methods of use in screening for compounds that
modulate sodium channel activity.

BACKGROUND OF THE INVENTION

Sodium channels are voltage-gated transmembrane proteins that are
15 involved in the generation of action potentials in electrically excitable cells such as
neurons and muscle cells. They are responsible for the cellular uptake of sodium
during electrical signals in cell membranes. The channels are members of a multigene
family of transmembrane proteins and are typically composed of a large
transmembrane pore-forming α -subunit and three smaller accessory β -subunits
20 (Cattrall et al., Adv Neurol 1999; 79:441-56). The primary structure of α -subunits is
conserved among different sub-types and species. The α -subunit is all that is required
for the channel to be fully functional, however, the β -subunits have been shown to
modulate the function of the channel. Specifically, co-expression of rat β 1, β 2, and
 β 3 subunits with the Na(v)1.2a α -subunits in the tsA-201 sub-clone of HEK293 cells
25 shifted sodium channel activation and inactivation to more positive membrane
potentials. The β 3 subunit alone caused increased persistent sodium currents. (Qu et
al., Mol Cell Neurosci 2001;18(5):570-80).

Previous studies have demonstrated numerous different types of α -subunits, which are categorized based on their sensitivity to tetrodotoxin (a toxin produced by the puffer or fugu fish). Subunits that are inhibited by nanomolar concentrations of tetrodotoxin are generally referred to as tetrodotoxin-sensitive channels (TTX-S), while those that require at least micromolar concentrations for inhibition are referred to as tetrodotoxin-resistant channels (TTX-R).

Rapid entry of sodium ions into cells causes depolarization and generation of the action potential. Such entry of sodium ions through sodium channels in response to a voltage change on the plasma membrane in excitable cells plays a functional role in control of neuronal excitability in the central nervous system (CNS) and peripheral nervous system (PNS).

An increase in the rate of spontaneous firing in neurons is often observed in peripheral sensory ganglia following nerve injury (Ochoa and Torebjork, Brain 1980; 103(4):835-53.; Nordin et al., Pain 1984; 20(3):231-45; Matzner et al., J Neurophysiol 1994; 72(1):349-59; Woolf, Drugs 1994; 47 Suppl 5:1-9; discussion 46-7). It has been suggested that this hyperexcitability in neurons is due to altered sodium channel expression in some chronic pain syndromes (Tanaka et al., Neuroreport 1998; 9(6):967-72). Increased numbers of sodium channels leading to inappropriate, repetitive firing of the neurons have been reported in the tips of injured axons in various peripheral nervous tissues such as the DRG, which relay signals from the peripheral receptors to the central nervous system (Waxman and Brill, Biophys J 1978; 21(2):147-60; Devor et al., Neurosci Lett 1989; 102(2-3):149-54; Matzner and Devor, Brain Res 1992; 597(1):92-98). Transcripts encoding the α III subunit, which are present at only very low levels in control DRG neurons, are expressed at moderate to high levels in axotomized DRG neurons together with elevated levels of α I and α II mRNAs (Waxman et al, Brain Res Mol Brain Res 1994; 22(1-4):275-89). Conversely, transcripts of sodium channel α subunits in the sensory nervous system are down-regulated in DRG neurons following axotomy (Dib-Hajj et al., Proc Natl Acad Sci U S A. 1996; 93(25):14950-4). Furthermore, the partial efficacy of sodium blocking agents is well documented in patients treated for neuropathic pain (Omana-Zapata et al., Pain 1997; 72(1-2):41-9; Rizzo, J Neurophysiol 1997; 77(1):236-46), providing an important link between increased sodium channel expression and

neuropathic pain. Therefore, alterations in sodium channel expression and subsequent function may be a key molecular event underlying the pathophysiology of pain after peripheral nerve injury.

5 The partial type III isoform (α -subunit) of the human sodium channel gene, SCN3A, isolated from placenta, was first described by Malo et al. (Proc Natl Acad Sci U S A 1994; 91(8):2975-9; GenBank Accession No. S69887). Two alternative isoforms, neonatal and adult forms, of SCN3A were thereafter identified in human brain tissue by Lu and Brown (J Mol Neurosci 1998;10(1):67-70; GenBank Accession Nos. AF035685 and AF035686, respectively). These isoforms contained a
10 92 amino acid insert within a region containing putative splice sites (identified through sequence homology with the rat type III brain sequence). The complete coding sequences for human SCN3A genomic DNA and mRNA (and the corresponding protein sequence) also cloned from human brain, was described by Clare et al. (Ann NY Acad Sci. 1999;868:80-3; GenBank Accession Nos. AJ251507
15 (SEQ ID NO: 3-Figure 3) and AF225987 (SEQ ID NO: 4-Figure 4, respectively).

Most recently, in 2000, Jeong et al. submitted to GenBank an mRNA sequence encoding a splice variant of human SCN3A (Accession No. AF225987; SEQ ID NO: 5-Figure 5). The amino acid sequence of this splice variant contained a
20 49-amino acid insert from residues 624 to 673 (SEQ ID NO: 6 - Figure 6), when compared with the sequence described by Clare et al. (*supra*).

There remains a need in the art to identify and characterize additional human sodium channels and variants thereof, in order to assist in the identification of drug candidates that can be used to treat conditions involving or associated with over- or under-expression, or over- or under-activated sodium channels.

25

SUMMARY OF THE INVENTION

The present invention provides a novel splice variant of human sodium channel III α subunit, designated herein as "hNaIII18", having the amino acid sequence of SEQ ID NO: 2 (Figure 2).

30 The present application also provides an isolated nucleic acid having a nucleotide sequence encoding the amino acid sequence of SEQ ID NO: 2. In one embodiment, the nucleic acid has the nucleotide sequence of SEQ ID NO: 1 (Figure

1). In another embodiment, the nucleic acid has a nucleotide sequence that is a degenerate variant of SEQ ID NO: 1. In yet another embodiment, the invention provides an isolated nucleic acid that hybridizes under stringent conditions to a nucleic acid having the nucleotide sequence of SEQ ID NO: 1, and preferably encodes
5 a protein having the same function as a protein having the amino acid sequence of SEQ ID NO: 2.

The isolated nucleic acid encoding hNaIII18 can be a part of a recombinant vector, *e.g.*, for cloning, expression, and/or expansion. An expression vector comprises the nucleic acid encoding hNaIII18 operably associated with an
10 expression control sequence. The invention further provides host cells containing such a vector, and methods for producing the hNaIII18 subunit polypeptide using such host cells.

In addition, the invention provides an isolated nucleic acid oligonucleotide, such as a primer or probe, of at least 10 bases, more particularly of at
15 least 20, and more particularly of at least 30 bases, which oligonucleotide has a nucleotide sequence identical to a corresponding nucleotide sequence of the same number of contiguous bases in SEQ ID NO: 1, or its complement, which nucleotide sequence is unique and specific to the nucleotide sequence of SEQ ID NO: 1, and/or different from corresponding oligonucleotide sequences encoding known sodium
20 channel subunits. The invention also provides an antibody that preferentially binds the hNaIII18 subunit protein of the invention compared to other known sodium channel subunits.

The present invention further provides a method for detecting expression of hNaIII18 in a cell or sample derived from a cell, which method
25 comprises: (i) detecting mRNA encoding hNaIII18 in a cell or in a sample derived from a cell suspected of expressing hNaIII18; or (ii) detecting hNaIII18 protein in a cell or in a sample derived from a cell with an antibody of the invention.

The present invention further provides an assay system for identifying modulators of hNaIII18 subunit containing sodium channels. The assay system
30 comprises at least one cell genetically engineered to express or overexpress hNaIII18 as part of a functional sodium channel, which can be used to screen for and thereby identify modulators of a hNaIII18-containing sodium channel. In a preferred

embodiment, cells useful in conducting the assay are mammalian cells useful in such screening methods including, *e.g.*, human embryonic kidney cells such as HEK293 cells, or cells such as *Xenopus* cells

BRIEF DESCRIPTION OF THE DRAWINGS

Figure 1 shows the cDNA sequence of hNaIII18 of the present invention.

Figure 2 shows the amino acid sequence of hNaIII18 of the present invention.

Figure 3 shows the cDNA sequence of human SCN3A of Clare et al. (*supra*) (GenBank Accession No. AJ251507).

Figure 4 shows the amino sequence of human SCN3A of Clare et al. (*supra*) (GenBank Accession No. AJ251507).

Figure 5 shows the cDNA of a human sodium channel α -subunit variant by Jeong et al. (GenBank Accession No. AF225987).

Figure 6 shows the amino acid sequence a human sodium channel α -subunit variant by Jeong et al. (GenBank Accession No. AF225987).

Figure 7 shows a cDNA alignment of the hNaIII18 of the present invention, with that of the human SCN3A of Clare et al. (*supra*), and that of Jeong et al. (*supra*)

Figure 8 shows the amino acid alignment of the hNaIII18 of the present invention, with that of the human SCN3A of Clare et al. (*supra*), and that of Jeong et al. (*supra*)

Figure 9A-D shows results of electrophysiology of hNaIII18-transfected HEK293 cells. Figure 9A demonstrates the activation threshold voltage; Figure 9B, the steady state $V_{1/2}$ inactivation voltage; Figure 9C, the recovery time after inactivation; and Figure 9D, the inactivation kinetics.

DETAILED DESCRIPTION OF THE INVENTION

The present invention is based, in part, on the discovery of a splice variant of the human NaIII channel α subunit. The human NaIII α subunit isoform, designated herein as "hNaIII18", was cloned by RT-PCR from human embryonic

brain total RNA (Clontech, Palo Alto, CA), using human NaIII specific primers. Primers were designed from a sequence identified by searching the NCBI Human Genome database, using the human NaIII mRNA sequence (GenBank accession no. AJ251507) using reverse-transcriptase PCR (RT-PCR). PCR fragments were cloned
5 into the mammalian expression vector and the complete DNA sequence was determined.

The hNaIII18 sequence of the invention contains an additional 147 nucleotides that do not appear in the human NaIII cDNA mentioned above (SEQ ID NO: 3). Splicing in this region (nucleotides +9 to +96) had been described for the rat
10 NaIII sodium channel, but not for the human NaIII channel when this work was initiated. The nucleotide sequence of Jeong et al. 2000, *supra*, also containing the 147 nucleotide insert and encoding an amino acid sequence similar to that of SEQ ID NO: 2, was deposited in GenBank (Accession No. AF225987, SEQ ID NO: 5), and is described in International PCT publication WO 01/96552 (in Japanese). The novel
15 sequence (SEQ ID NO: 1) presented herein differs from that of SEQ ID NO:5 by 37 nucleotides out of 6093 aligned. None of the differences are found within the 147-nucleotide insertion. The amino acid sequence presented herein in SEQ ID NO: 2, differs from the SEQ ID NO:5 amino acid sequence by 12 amino acids out of 2000, with none of the differences being found in the region containing the 49 amino acid
20 insert.

Transient transfection of the novel splice variant of the invention (SEQ ID NO: 1) results in expression of functional sodium channels in mammalian cells (cell line HEK293). Stable transfection and expression of the hNaIII18 also was achieved in HEK293 cells.

25 Protein expression was confirmed in the stably transfected HEK293 cells by immunocytochemistry and Western blotting. A protein having a size of about 220 kD protein, corresponding to the expected molecular weight of hNaIII18 was identified. Functional hNaIII18 activity was confirmed by electrophysiology.

Thus, the present invention advantageously provides hNaIII18 protein,
30 including fragments and derivatives thereof; hNaIII18-encoding nucleic acids, and portions thereof including oligonucleotide primers and probes surrounding and within the region containing the 147 nucleotide insert, and hNaIII18 regulatory sequences;

hNaIII18-specific antibodies; and related methods of using these materials to detect the presence of hNaIII18 proteins or nucleic acids.

The present invention also provides an assay method for screening to identify selective modulators of hNaIII18-containing sodium ion channel activity.

5 The method involves detecting whether a test compound increases or decreases the activity of the sodium channel, as determined, *e.g.*, by measuring current phase (electrophysiology) and ion selectivity. The assay method is preferably conducted using at least one host cell that expresses or overexpresses a functional sodium channel comprising hNaIII18, or a membrane preparation prepared therefrom. In one
10 embodiment, the test compound inhibits (antagonizes) the activity of the sodium channel. In another embodiment, the test compound potentiates (agonizes) the activity of the sodium channel. The test system preferably involves the use of an appropriate cell culture medium to permit cell growth and viability, as well as tissue culture plates or arrays containing the host cells in the cell culture medium. In
15 specific embodiments, host cells are mammalian cell lines such as, *e.g.*, the HEK293 cell line, although appropriate cells from other organisms, such as, *e.g.*, *Xenopus* cells, can alternatively be utilized.

The specification and figures include the following nucleotide or amino acid sequences: hNaIII18 polynucleotide (SEQ ID NO:1); hNaIII18 amino acid
20 sequence (SEQ ID NO:2); SCN3A nucleotide sequence (SEQ ID NO:3; Clare et al., *supra*; GenBank AJ251507); SCN3A amino acid sequence (SEQ ID NO:4; Clare et al., *supra*; GenBank AJ201507); SCN3A splice variant nucleotide sequence (SEQ ID NO:5; Jeong et al., *supra*; GenBank AF225987); SCN3A splice variant amino acid
25 sequence (SEQ ID NO:6; Jeong et al., *supra*; GenBank AF225987); forward primer utilized in Example 1 (SEQ ID NO:7); and reverse primer utilized in Example 1 (SEQ ID NO:8).

General Definitions

30 The following definitions are provided for clarity and illustrative purposes only, and are not intended to limit the scope of the invention.

As used herein, the term "isolated" means that the referenced material is removed from the environment in which it is normally found. Thus, an isolated

biological material can be free of cellular components, *i.e.*, components of the cells in which the material is found or produced in nature. In the case of nucleic acid molecules, an isolated nucleic acid includes a PCR product, an mRNA, a cDNA, or a restriction fragment. In another embodiment, an isolated nucleic acid is preferably excised from the chromosome in which it may be found, and more preferably is no longer joined to non-regulatory, non-coding regions, or to other genes, located upstream or downstream of the gene contained by the isolated nucleic acid molecule when found in the chromosome. In yet another embodiment, the isolated nucleic acid lacks one or more naturally occurring introns. Isolated nucleic acid molecules include sequences inserted into plasmids, cosmids, artificial chromosomes, phages and the like. Thus, in a specific embodiment, a recombinant nucleic acid is an isolated nucleic acid. An isolated protein may be associated with other proteins or nucleic acids, or both, with which it associates in the cell, or with cellular membranes if it is a membrane-associated protein. A protein expressed from a vector in a cell, particularly a cell in which the protein is normally not expressed, is also regarded as isolated. An isolated organelle, cell, or tissue is removed from the anatomical site in which it is found in a cell or an organism. An isolated material may be, but need not be, purified. As used herein to refer to nucleic acids, the term "isolated" does not encompass man-made genomic or cDNA libraries.

The term "purified" as used herein refers to material that has been isolated under conditions that reduce or eliminate the presence of unrelated materials, *i.e.*, contaminants, including native materials from which the material is obtained. For example, a purified protein is preferably substantially free of other proteins or nucleic acids with which it is associated in a cell; a purified nucleic acid molecule is preferably substantially free of proteins or other unrelated nucleic acid molecules with which it can be found within a cell. As used herein, the term "substantially free" is used operationally, in the context of analytical testing of the material. Preferably, purified material substantially free of contaminants. Purity can be evaluated by chromatography, gel electrophoresis, immunoassay, composition analysis, biological assay, and other methods known in the art.

Methods for purification are well-known in the art. For example, nucleic acids can be purified by precipitation, chromatography (including preparative

solid phase chromatography, oligonucleotide hybridization, and triple helix chromatography), ultracentrifugation, and other means. Polypeptides and proteins can be purified by various methods including, without limitation, preparative disc-gel electrophoresis, isoelectric focusing, HPLC, reversed-phase HPLC, gel filtration, ion exchange and partition chromatography, precipitation and salting-out chromatography, extraction, and countercurrent distribution. For some purposes, it is preferable to produce the protein in a recombinant system so that it contains an additional sequence tag that facilitates purification, such as, but not limited to, a polyhistidine sequence (His®-tag; Novagen, Madison, WI), or a sequence that specifically binds to an antibody, such as the FLAG® tag (Sigma, St. Louis, MO), HA-tag (Roche Diagnostics, Indianapolis, IN), or that can be column-purified such as via the use of glutathione-S-transferase (GST). The polypeptide can then be purified from a crude lysate of the host cell by chromatography on an appropriate solid-phase matrix. Alternatively, antibodies produced against the protein or against peptides derived therefrom can be used as purification reagents. Cells can be purified by various techniques, including centrifugation, matrix separation (*e.g.*, nylon wool separation), panning and other immunoselection techniques, depletion (*e.g.*, complement depletion of contaminating cells), and cell sorting (*e.g.*, fluorescence activated cell sorting (FACS)). Other purification methods are possible. A purified material may contain less than about 50%, preferably less than about 75%, and most preferably less than about 90%, by weight of the cellular components with which it was originally associated. The "substantially pure" indicates the highest degree of purity that can be achieved using conventional purification techniques known in the art.

In a specific embodiment, the term "about" or "approximately" means plus or minus 10% of the stated numerical value or range.

As use herein, the term "ion channel" refers to a transmembrane pore that presents a hydrophilic channel for ions to cross a lipid bilayer down their electrochemical gradients. In a preferred embodiment, the ion channel is a voltage-gated sodium ion channel. A "sodium channel" is an ion channel that is selective for sodium ions.

A "sample" as used herein refers to a biological material that can be obtained and tested for the presence or expression of: (i) an hNaIII18 subunit-containing ion channel; or (ii) an hNaIII18 subunit protein; or (iii) an hNaIII18 subunit-encoding nucleic acid. Such samples can be obtained from animal, preferably mammalian, and more preferably human subjects, and include tissue samples, especially CNS or PNS tissues, as well as cell cultures derived from such tissues. Alternatively, such samples can comprise cells genetically engineered to express or overexpress an hNaIII18 subunit-containing ion channel or an hNaIII18 subunit protein. Such cells are preferably eukaryotic, but may alternatively be prokaryotic cells. Eukaryotic cells are preferably mammalian cells, but may alternatively be *Xenopus* cells.

Non-human animals include, without limitation, laboratory animals such as mice, rats, rabbits, hamsters, guinea pigs, etc.; domestic animals such as dogs and cats; and farm animals such as sheep, goats, pigs, horses, and cows.

The term "modulator" refers to a compound that binds to an ion channel comprising the hNaIII18 subunit protein of the invention and differentially affects the activity of the ion channel in response to a stimulus that normally activates the function of that ion channel when compared to the activity of the ion channel not contacted with the compound. Ion channel activity can be measured, *e.g.*, using electrophysiological techniques, or according to other known methods in the art. In a preferred embodiment, the ion channel is a sodium channel.

The terms "inhibitor" and antagonist refer to a compound that binds to the ion channel comprising hNaIII18, and blocks, inhibits, impedes or reduces the activity of that ion channel.

An "agonist" is defined as a compound that binds to the ion channel comprising hNaIII18, and promotes, enhances, stimulates or potentiates the normal biological function of the sodium channel. A "partial agonist" binds as to the ion channel or a subunit thereof, as does a full agonist, but promotes only partial function.

As used herein the term "transfected cell" or "transformed cell" refers to a host cell that has been genetically engineered to express or overexpress a nucleic acid encoding a hNaIII18 subunit, preferably in combination with one or more β subunits such as, *e.g.*, β -subunits 1-3 as described in GenBank Accession Nos.

U87445, AF007783, AH005825, AF007783, AF04948, L10338 and L16242, among others. Any cell can be used, preferably a eukaryotic cell, and more preferably a vertebrate cells, preferably a mammalian cell, or a *Xenopus* cell. Such cells additionally can be genetically engineered to coexpress or overexpress a different sodium channel subunit. Such genetically engineered cells include those cells into which one or more heterologous hNaIII18-encoding nucleic acids have been introduced and are expressed or overexpressed. Such genetically engineered cells also include those cells engineered to express or overexpress one or more endogenous hNaIII18 subunits, for example, by gene activation technology.

Such cells are particularly suitable to conduct an assay to screen for compounds that modulate the function of the hNaIII18 subunit-containing sodium channel in response to an appropriate stimulus (*e.g.*, TTX). An "assay method" typically makes use of one or more such cells, *e.g.*, in a microwell plate or some other culture system. The effects of a test compound can be determined on a single cell or on a collection of cells sufficient to allow measurement of ionic current, activation threshold, or ionic permeability characteristics of the hNaIII18 subunit-containing sodium channels. For example, single cells can be tested, *e.g.*, by use of patch clamp or other appropriate electrophysiological techniques.

A "test compound" or "candidate compound" is any molecule that can be tested for its ability to bind to the hNaIII18 subunit-containing sodium channel, or to a subunit thereof, and preferably modulate on the activity of the hNaIII18 subunit-containing sodium channel. A compound that binds and modulates a hNaIII18 subunit-containing sodium channel is a "lead compound" suitable for further testing and development.

The term "ligand" can alternatively be used to refer to any compound or peptide or polypeptide that binds to and modulates the activity of a hNaIII18 subunit, or a sodium channel comprising hNaIII18.

The term "pain disorder" includes chronic pain, defined as pain lasting longer than one month (Bonica, *Semin Anesth* 1986, 5:82-99), and is characterized by unrelenting persistent pain that is not amenable to routine pain control methods. The term "pain disorder" also includes neuropathic pain and nociceptive pain.

“Chronic pain” can be defined as pain lasting longer than one month (Bonica, *Semin Anesth* 1986, 5:82-99), and is characterized by unrelenting persistent pain that is not amenable to routine pain control methods. Chronic pain includes, but is not limited to, inflammatory pain, postoperative pain, cancer pain, osteoarthritis pain associated with metastatic cancer, trigeminal neuralgia, acute herpetic and postherpetic neuralgia, diabetetic neuropathy, causalgia, brachial plexus avulsion, occipital neuralgia, reflex sympathetic dystrophy, fibromyalgia, gout, phantom limb pain, burn pain, pain associated with spinal cord injury, multiple sclerosis, reflex sympathetic dystrophy and lower back pain and other forms of neuralgia, neuropathic, and idiopathic pain syndromes.

“Neuropathic pain” can be caused by injury or infection of peripheral sensory nerves. It includes, but is not limited to pain from peripheral nerve trauma, herpes virus infection, diabetes mellitus, causalgia, plexus avulsion, neuroma, limb amputation, and vasculitis. Neuropathic pain is also caused by nerve damage from chronic alcoholism, human immunodeficiency virus infection, hypothyroidism, uremia, or vitamin deficiencies. Neuropathic pain includes but is not limited to pain caused by nerve injury such as, for example, the pain from which diabetics suffer.

Chronic and neuropathic types of pain generally arises from injury to the peripheral or central nervous tissue.

“Nociceptive pain” is due to activation of pain-sensitive nerve fibers, either somatic or visceral. Nociceptive pain generally results as a response to direct tissue damage. The initial trauma causes the release of several chemicals including bradykinin, serotonin, substance P, histamine, and prostaglandin. When somatic nerves are involved, the pain is typically experienced as aching or pressure-like.

Molecular Biology Definitions

In accordance with the present invention there may be employed conventional molecular biology, microbiology, and recombinant DNA techniques within the skill of the art. See, *e.g.*, Sambrook, Fritsch & Maniatis, *Molecular Cloning: A Laboratory Manual*, Second Edition (1989) Cold Spring Harbor Laboratory Press, Cold Spring Harbor, New York (herein "Sambrook et al., 1989"); *DNA Cloning: A Practical Approach*, Volumes I and II (D.N. Glover ed. 1985);

Oligonucleotide Synthesis (M.J. Gait ed. 1984); Nucleic Acid Hybridization [B.D. Hames & S.J. Higgins eds. (1985)]; Transcription And Translation [B.D. Hames & S.J. Higgins, eds. (1984)]; Animal Cell Culture [R.I. Freshney, ed. (1986)]; Immobilized Cells And Enzymes [IRL Press, (1986)]; B.Perbal, A Practical Guide To Molecular Cloning (1984); F.M. Ausubel et al. (eds.), Current Protocols in Molecular Biology, John Wiley & Sons, Inc. (1994).

"Amplification" of DNA as used herein denotes the use of exponential amplification, techniques such as polymerase chain reaction (PCR), and non-exponential amplification, such as linked linear amplification, to increase the concentration of a particular DNA sequence within a mixture of DNA sequences. For a description of PCR see Saiki et al., Science 1988, 239:487. For a description of linked linear amplification, see U.S. Patent Nos. 6,335,184 and 6,027,923 and Reyes et al. Clinical Chemistry 2001; 47: 131-40; Wu et al. Genomics 1989; 4: 560-569.

As used herein, "sequence-specific oligonucleotides" refers to related sets of oligonucleotides that can be used to detect allelic variations or mutations in the hNAIII18 gene, or can be used for amplification of an hNAIII18 encoding-nucleic acid.

The nucleic acid molecules (polynucleotides) described herein may be flanked by natural regulatory (expression control) sequences, or may be associated with heterologous sequences, including promoters, internal ribosome entry sites (IRES) and other ribosome binding site sequences, enhancers, response elements, suppressors, signal sequences, polyadenylation sequences, introns, 5'- and 3'- non-coding regions, and the like. The nucleic acid molecules may also be modified by many means known in the art. Non-limiting examples of such modifications include methylation, "caps", substitution of one or more of the naturally occurring nucleotides with an analog, and internucleotide modifications such as, for example, replacement with uncharged linkages (*e.g.*, methyl phosphonates, phosphotriesters, phosphoroamidates, carbamates, etc.) and with charged linkages (*e.g.*, phosphorothioates, phosphorodithioates, etc.). Polynucleotides may contain one or more additional covalently linked moieties, such as, for example, proteins (*e.g.*, nucleases, toxins, antibodies, signal peptides, poly-L-lysine, etc.), intercalators (*e.g.*, acridine, psoralen, etc.), chelators (*e.g.*, metals, radioactive metals, iron, oxidative

metals, etc.), and alkylators. The polynucleotides may be derivatized by formation of a methyl or ethyl phosphotriester or an alkyl phosphoramidate linkage. Furthermore, the polynucleotides herein may also be modified with a label capable of providing a detectable signal, either directly or indirectly. Exemplary labels include

5 radioisotopes, fluorescent molecules, biotin, and the like.

A "coding sequence" or a sequence "encoding" an expression product, such as an RNA, polypeptide, protein, or enzyme, is a nucleotide sequence that, when expressed, results in the production of that RNA or polypeptide, *i.e.*, the nucleotide sequence encodes an amino acid sequence for that polypeptide. A coding sequence or

10 "open reading frame (ORF)" for a polypeptide will typically include a start codon (usually ATG) and a stop codon.

The term "gene", also called a "structural gene" refers to a basic unit of hereditary material. Specifically a gene is an ordered sequence of DNA nucleotide bases that encodes one polypeptide chain (via mRNA). The gene includes regions

15 preceding and following the coding region (such as promoter sequences, a 5'-untranslated region, and a 3'-untranslated region, which affect, for example, the conditions under which the gene is expressed) as well as (in eukaryotes) intervening sequences (introns) between individual coding segments (exons).

A "promoter sequence" is a DNA regulatory region capable of binding

20 RNA polymerase in a cell and initiating transcription of a downstream (3' direction) coding sequence. For purposes of defining the present invention, the promoter sequence is bounded at its 3' terminus by the transcription initiation site and extends upstream (5' direction) to include the minimum number of bases or elements necessary to initiate transcription at levels detectable above background. Within the

25 promoter sequence will be found a transcription initiation site (conveniently defined for example, by mapping with nuclease S1), as well as protein binding domains (consensus sequences) responsible for the binding of RNA polymerase. The present invention includes the hNaIII18 gene promoter found in the genome, which can be operatively associated with a hNaIII18 coding sequence with a heterologous coding

30 sequence.

The term "host cell" means any cell of any organism that is selected, modified, transformed, grown, or used or manipulated in any way, for the production

of a substance by the cell, for example, the expression by the cell of a gene, a DNA or RNA sequence, or a polypeptide. Host cells can further be used for screening or other assays, as described *infra*.

5 A coding sequence is "under the control of" or "operatively associated with" transcriptional and translational control sequences in a cell when such control sequences operate to effect RNA polymerase transcription of the coding sequence into mRNA, which is then trans-RNA spliced (if it contains introns) and translated, in the case of mRNA, into the protein encoded by the coding sequence.

10 The terms "express" and "expression" mean allowing or causing the information in a gene or cDNA or mRNA sequence to become manifest, for example, by producing a protein by activating the cellular functions involved in transcription and translation of a corresponding gene, cDNA or mRNA sequence. A gene or cDNA sequence is expressed in or by a cell to form an "expression product" such as a protein. The expression product itself, *e.g.*, the resulting protein, may also be said to
15 be "expressed" by the cell. An expression product can be characterized as intracellular, extracellular, transmembrane, or secreted depending on the particular product. The hNaIII18 subunit protein of the invention is typically expressed as a transmembrane protein with intracellular and extracellular domains.

The term "transfection" means the introduction of a "foreign" (*i.e.*,
20 extrinsic or extracellular) gene, DNA or RNA sequence into a host cell so that the host cell will express the introduced gene or sequence to produce a desired substance, typically a protein encoded by the introduced gene or sequence. The introduced gene or sequence may also be called a "cloned" or "foreign" or "heterologous" gene or sequence, and may include regulatory or control sequences, such as start, stop,
25 promoter, signal, secretion, or other sequences used by a cell's genetic machinery. The gene or sequence may include non-functional sequences or sequences with no known function.

The term "transformation" refers to the process by which DNA is introduced from the surrounding medium into a prokaryotic host cell.

30 The term "transduction" refers to the introduction of DNA into a prokaryotic host cell via a bacterial virus, or bacteriophage.

A prokaryotic or eukaryotic host cell that receives and expresses introduced DNA or RNA has been "transformed" and is a "transformant" or a "clone." The DNA or RNA introduced into a host cell can come from any source, including cells of the same genus or species as the host cell, or cells of a different genus or species, or synthetic sequences.

The transformed cells of the invention are particularly suitable for an assay system for the detection of compounds that modulate the function of hNaIII18 subunit-containing sodium channels in response to activation, *e.g.*, in response to exposure TTX. An "assay method" makes use of one or more such cells, *e.g.*, in a microwell plate or some other culture or assay system to permit evaluation of the effects of a test compound on the cell(s), *e.g.*, by measuring ionic current or activation threshold characteristics of the hNaIII18 subunit-containing sodium channel.

The term "recombinantly engineered cell" refers to any prokaryotic or eukaryotic cell that has been manipulated to express or overexpress the hNaIII18 subunit by any appropriate method, including transfection, transformation or transduction. This term also includes endogenous activation of a hNaIII18 gene in a cell that does not normally express hNaIII18 or that expresses the protein at a sub-optimal level.

The terms "vector", "cloning vector" and "expression vector" mean the vehicle by which a DNA or RNA sequence (*e.g.*, a foreign gene) can be introduced into a host cell, so as to transform the host and promote expression (*e.g.*, transcription and translation) of the introduced sequence. Vectors include plasmids, cosmids, phages, viruses, etc.; they are discussed in greater detail below.

Vectors typically comprise the DNA of a transmissible agent, into which foreign DNA is inserted. A common way to insert one segment of DNA into another segment of DNA involves the use of restriction enzymes to cleave DNA at specific restriction sites. A "cassette" refers to a DNA coding sequence or segment of DNA that codes for an expression product that can be inserted into a vector at defined restriction sites. The cassette restriction sites are designed to ensure insertion of the cassette in the proper reading frame. Generally, foreign DNA is inserted at one or more restriction sites of the vector DNA, and then is carried by the vector into a host cell along with the transmissible vector DNA. A segment or sequence of DNA

having inserted or added DNA, such as an expression vector, can also be called a "DNA construct." A common type of vector is a plasmid. A plasmid vector often contains coding DNA and promoter DNA and has one or more restriction sites suitable for inserting foreign DNA. Coding DNA is a DNA sequence that encodes a particular amino acid sequence for a particular protein. Promoter DNA is a DNA sequence that initiates, regulates, or otherwise mediates or controls the expression of the coding DNA. Promoter DNA and coding DNA may be from the same gene or from different genes, and may be from the same or different organisms. A large number of vectors, including plasmid and fungal vectors, have been described for replication and/or expression in a variety of eukaryotic and prokaryotic hosts. Non-limiting examples include pKK plasmids (Clontech), pUC plasmids, pET plasmids (Novagen, Inc., Madison, WI), pRSET or pREP plasmids (Invitrogen, San Diego, CA), or pMAL plasmids (New England Biolabs, Beverly, MA), and many appropriate host cells. Recombinant cloning vectors will often include one or more replication systems for cloning or expression, one or more markers for selection in the host, *e.g.*, antibiotic resistance, and one or more expression cassettes.

The term "expression system" means a host cell and compatible vector under suitable conditions, *e.g.*, for the expression of a protein coded for by foreign DNA carried by the vector and introduced to the host cell. Common expression systems include *E. coli* host cells and plasmid vectors, insect host cells and baculovirus vectors, and mammalian host cells and vectors.

The term "heterologous" refers to a combination of elements not naturally occurring. For example, heterologous DNA refers to DNA not naturally present in that cell. Alternatively, heterologous DNA refers to combinations of sequences that do not naturally occur together in that cell, *e.g.*, promoter sequences from a gene from one cell type linked to coding sequences of a gene that is not normally controlled by that promoter or expressed by another cell type. Preferably, the heterologous DNA includes a gene foreign to the cell. A heterologous expression regulatory element is such an element operatively associated with a different gene than the one it is operatively associated with in nature. In the context of the present invention, a hNaIII18 gene is heterologous to the vector DNA in which it is inserted

for cloning or expression purposes, and is heterologous to a host cell containing such a vector in which it is expressed, *e.g.*, a HEK cell.

The terms "mutant" and "mutation" mean any detectable change in genetic material, *e.g.*, DNA, or any process, mechanism, or result of such a change.

5 This includes gene mutations in which the structure (*e.g.*, DNA sequence) of a gene is altered; any gene or DNA arising from any mutation process; and any expression product (*e.g.*, protein or enzyme) expressed by a non-silent modification of a gene or DNA sequence. The term "variant" may also be used to indicate a modified or altered gene, DNA sequence, polypeptide, cell, etc., *i.e.*, any kind of mutant therefrom.

10 "Sequence-conservative variants" or "degenerate variants" of a polynucleotide sequence are those in which a change of one or more nucleotides in a given codon position results in no alteration in the amino acid encoded at that position.

"Function-conservative variants" are those in which a given amino acid
15 residue in a protein has been changed without substantially altering the function of the polypeptide, including, but not limited to, replacement of an amino acid with a residue having similar properties (such as, for example, polarity, hydrogen bonding potential, acidic, basic, hydrophobic, aromatic, and the like). Amino acids with similar properties are well known in the art. For example, arginine, histidine and lysine are
20 hydrophilic-basic amino acids and may be interchangeable. Similarly, isoleucine, a hydrophobic amino acid, may be replaced with leucine, methionine or valine. Such changes are expected to have little or no effect on the apparent molecular weight, isoelectric point, or function of the protein. Amino acid residues may be varied in a protein so that the percent amino acid sequence identity between the original protein
25 and the variant may be, for example, at least 70%, 80%, 90%, 95% or 99%, as determined according to a default alignment scheme such as by the Cluster Method, wherein similarity is based on the MEGALIGN algorithm, or BLAST. A "function-conservative variant" of the present invention includes those polypeptides having the above-described amino acid sequence identities, and having the same or substantially
30 similar functions as the native or parent hNaIII18 subunit protein of the invention

As used herein, the term "homologous" refers to the relationship between proteins that possess a "common evolutionary origin," including proteins

from superfamilies (*e.g.*, the immunoglobulin superfamily) and homologous proteins from different species (*e.g.*, myosin light chain, etc.) (Reeck et al., Cell 1987, 50:667). Such proteins (and their encoding genes) have sequence homology, as reflected by their sequence similarity or sequence identity, whether in terms of percent similarity or the presence of specific residues or motifs at conserved positions.

Accordingly, the term "sequence similarity" or "sequence identity" refers to the degree of identity or correspondence between nucleic acid or amino acid sequences of proteins that may or may not share a common evolutionary origin (see Reeck et al., *supra*). However, in common usage and in the instant application, the term "homologous," when modified with an adverb such as "highly," may refer to sequence similarity and may or may not relate to a common evolutionary origin.

In a specific embodiment, two DNA sequences are "substantially homologous" or "substantially similar" when at least about 80%, and most preferably at least about 90, 95% or 99% of the nucleotides match over the defined length of the DNA sequences, as determined by sequence comparison algorithms, such as BLAST, FASTA, DNA Strider, etc. An example of such a sequence is an allelic or species variant of the specific hNaIII18 gene of the invention. Sequences that are substantially homologous can be identified by comparing the sequences using standard software available in sequence data banks, or in a Southern hybridization experiment under, for example, stringent conditions as defined for that particular system.

Similarly, in a particular embodiment, two amino acid sequences are "substantially homologous" or "substantially similar" when greater than 80%, 90%, 95% or 99% of the amino acids are identical. Preferably, the similar or homologous sequences are identified by alignment using, for example, the GCG (Genetics Computer Group, Program Manual for the GCG Package, Version 7, Madison, Wisconsin) pileup program, or any of the programs described above (BLAST, FASTA, etc.).

A nucleic acid molecule is "hybridizable" to another nucleic acid molecule, such as a cDNA, genomic DNA, or RNA, when a single stranded form of the nucleic acid molecule can anneal to the other nucleic acid molecule or its complement under the appropriate conditions of temperature and solution ionic

strength (see Sambrook *et al.*, *supra*). The conditions of temperature and ionic strength determine the "stringency" of the hybridization. For preliminary screening for homologous nucleic acids, low stringency hybridization conditions, using a T_m (melting temperature) in the range of about 55 °C with low salt and/or denaturant concentrations, can be used, *e.g.*, 5x SSC, 0.1% SDS, 0.25% milk, and no formamide; or 30% formamide, 5x SSC, 0.5% SDS. Moderate stringency hybridization conditions correspond to use of a higher T_m , and higher concentrations of salt and/or denaturants, *e.g.*, 40% formamide, with 5x or 6x SSC. High stringency hybridization conditions correspond to the highest T_m and concentrations of salt/and/or denaturants, *e.g.*, 68°C, 50% formamide, 5x or 6x SSC. SSC is a 0.15M NaCl, 0.015M Na-citrate buffer. Hybridization requires that the two nucleic acids contain complementary sequences, although depending on the stringency of the hybridization, mismatches between bases are possible. The appropriate stringency for hybridizing nucleic acids depends on the length of the nucleic acids and the degree of complementation, as known in the art. The greater the degree of similarity or homology between two nucleotide sequences, the higher the value of T_m for hybrids of nucleic acids having those sequences. The relative stability (corresponding to higher T_m) of nucleic acid hybridizations decreases in the following order: RNA:RNA, DNA:RNA, DNA:DNA. For hybrids of greater than 100 nucleotides in length, equations for calculating T_m have been derived (see Sambrook *et al.* 1989, *supra*, 9.50-9.51). For hybridization with shorter nucleic acids, *i.e.*, oligonucleotides, the position of mismatches becomes more important, and the length of the oligonucleotide determines its specificity (see Sambrook *et al.*, *supra*, 11.7-11.8). A minimum length for a hybridizable nucleic acid is at least about 10 nucleotides; preferably at least about 15 nucleotides; and more preferably at least about 20 nucleotides.

In a specific embodiment, the term "standard hybridization conditions" refers to a T_m of 55 °C, and utilizes conditions as set forth above. In a preferred embodiment, the T_m is about 60 °C; in a more preferred embodiment, the T_m is about 65 °C. In a specific embodiment, "high stringency" refers to hybridization and/or washing conditions at 68 °C, in 0.2 x SSC, at 42 °C in 50% formamide, 4x SSC, or under conditions that afford levels of hybridization equivalent to those observed under either of these two conditions.

As used herein, the term "oligonucleotide" refers to a nucleic acid, generally of at least 10, preferably at least 15, and more preferably at least 20 nucleotides, preferably no more than 100 nucleotides, that is hybridizable to a genomic DNA molecule, a cDNA molecule, or an mRNA molecule, or other nucleic acid of interest. Oligonucleotides can be labeled, *e.g.*, with $\gamma^{32}\text{P}$ -nucleotides or nucleotides to which a label, such as biotin, has been covalently conjugated. In one embodiment, a labeled oligonucleotide can be used as a probe to detect the presence of a nucleic acid. In another embodiment, oligonucleotides (one or both of which may be labeled) can be used as PCR primers, either for cloning a full length nucleic acid or a fragment of a nucleic acid encoding the hNaIII18 subunit, or to detect the presence of nucleic acids encoding hNaIII18. In a further embodiment, an oligonucleotide of the invention can form a triple helix with a hNaIII18-encoding DNA molecule. Generally, oligonucleotides are prepared synthetically, preferably on a nucleic acid synthesizer. Accordingly, oligonucleotides can be prepared with non-naturally occurring phosphoester analog bonds, such as thioester bonds, etc.

The present invention also provides antisense nucleic acids, which may be used to inhibit expression of the hNaIII18 subunit protein of the invention. Inhibition of hNaIII18 expression may be desired when upregulation of hNaIII18 expression or excessive activation of an hNaIII18-containing ion channel induces or otherwise contributes to an increase in pain or a pain disorder in a subject.

An "antisense nucleic acid" is a single stranded nucleic acid molecule, which may be DNA, RNA, a DNA-RNA chimera, or derivatives thereof, which, on hybridizing under cytoplasmic conditions with complementary bases in an RNA or DNA molecule, inhibits the expression or translation of the encoded gene. If the RNA is an mRNA transcript, the antisense nucleic acid is a counter-transcript or mRNA-interfering complementary nucleic acid. As presently used, "antisense" broadly includes RNA-RNA interactions, RNA-DNA interactions, and RNase-H mediated arrest. Antisense nucleic acid molecules can be encoded by a recombinant gene for expression in a cell (*e.g.*, U.S. Patent No. 5,814,500; U.S. Patent No. 5,811,234), or alternatively they can be prepared synthetically (*see, e.g.*, U.S. Patent No. 5,780,607).

In addition to antisense sequences, the present invention also provides ribozymes useful to inhibit hNaIII18 expression. Ribozyme technology is described further in Intracellular Ribozyme Applications: Principals and Protocols, Ed. Rossi and Couture, 1999, Horizon Scientific Press

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hNaIII18 Nucleic Acids

A polynucleotide molecule encoding hNaIII18, whether genomic DNA or cDNA, can be isolated from any source, particularly from a human cDNA or genomic library. Methods for obtaining specific polynucleotide molecules gene are well known in the art, as described above (see, *e.g.*, Sambrook *et al.*, 1989, *supra*). The DNA may be obtained by standard procedures known in the art from cloned DNA (*e.g.*, a DNA "library"), and preferably is obtained from a cDNA library prepared from tissues with high level expression of the encoded protein, by chemical synthesis, by cDNA cloning, or by the cloning of genomic DNA, or fragments thereof, purified from the desired cell (See, for example, Sambrook *et al.*, 1989, *supra*; Glover, D.M. (ed.), 1985, DNA Cloning: A Practical Approach, MRL Press, Ltd., Oxford, U.K. Vol. I, II). Clones derived from genomic DNA may contain regulatory and intron DNA regions in addition to coding regions. Clones derived from cDNA will not contain intron sequences. Whatever the source, the polynucleotide molecule should be cloned into a vector suitable for its propagation. Identification of a specific DNA fragment containing the desired hNaIII18-encoding sequence may be accomplished in a number of ways. For example, a portion of a hNaIII18 encoding polynucleotide molecule exemplified *infra* can be purified and labeled to prepare a labeled probe, and the generated DNA library may be screened by nucleic acid hybridization to the labeled probe (Benton and Davis, Science 1977, 196:180; Grunstein and Hogness, Proc. Natl. Acad. Sci. U.S.A. 1975, 72:3961). Those DNA fragments with substantial homology to the probe, such as an allelic variant from another individual, will hybridize. In a specific embodiment, highest stringency hybridization conditions are used to identify a homologous hNaIII18 gene.

Further selection can be carried out on the basis of the properties of the gene, *e.g.*, if the gene encodes a protein product having the same physicochemical profile (*i.e.*, isoelectric, electrophoretic, electrophysiological, amino acid composition,

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partial or complete amino acid sequence, antibody binding activity, or ligand binding profile) of the hNaIII18 subunit protein disclosed herein. Thus, the presence of the nucleic acid may be detected by assays based on the physical, chemical, immunological, or functional properties of its expressed product.

5 Other DNA sequences which encode substantially the same amino acid sequence as a hNaIII18 gene may be used in the practice of the present invention. These include but are not limited to allelic variants, species variants, sequence conservative variants, and function conservative variants.

10 Amino acid substitutions may also be introduced to substitute an amino acid with a particularly preferable property. For example, a Cys may be introduced at a potential site for disulfide bridges with another Cys.

Polynucleotide molecules encoding the hNaIII18 subunit, and the encoded polypeptide, derivatives and analogs thereof, can be produced by various methods known in the art. The manipulations which result in their production can occur at the gene or protein level. For example, the cloned hNaIII18 gene or cDNA sequence can be modified by any of numerous strategies known in the art (Sambrook *et al.*, 1989, *supra*). The sequence can be cleaved at appropriate sites with restriction endonuclease(s), followed by further enzymatic modification if desired, isolated, and ligated *in vitro*. In the production of the polynucleotide molecule encoding a derivative or analog of hNaIII18, care should be taken to ensure that the modified polynucleotide sequence remains within the same translational reading frame as the hNaIII18 gene, uninterrupted by premature translational stop signals.

20 Additionally, the encoding nucleic acid sequence can be mutated *in vitro* or *in vivo* to create and/or destroy translation, initiation, and/or termination sequences, or to create variations in coding regions and/or form new restriction endonuclease sites or destroy preexisting ones, to facilitate further *in vitro* modification. Such modifications can be made to introduce restriction sites and facilitate cloning the polynucleotide molecule into an expression vector. Any technique for mutagenesis known in the art can be used, including but not limited to, *in vitro* site-directed mutagenesis (Hutchinson, C., *et al.*, J. Biol. Chem. 1978; 253:6551; Zoller and Smith, DNA 1984; 3:479-488; Oliphant *et al.*, Gene 1986; 44:177; Hutchinson *et al.*, Proc. Natl. Acad. Sci. U.S.A. 1986; 83:710), use of TAB

linkers (Pharmacia), etc. PCR techniques are preferred for site directed mutagenesis (see Higuchi, 1989, "Using PCR to Engineer DNA", in PCR Technology: Principles and Applications for DNA Amplification, H. Erlich, ed., Stockton Press, Chapter 6, pp. 61-70).

5 The identified and isolated polynucleotide molecule can then be inserted into an appropriate cloning vector. A large number of vector-host systems known in the art may be used. Possible vectors include, but are not limited to, plasmids or modified viruses, but the vector system must be compatible with the host cell used. Examples of vectors include, but are not limited to, *E. coli*, bacteriophages
10 such as lambda derivatives, or plasmids such as Bluescript, pBR322 derivatives or pUC plasmid derivatives, *e.g.*, pGEX vectors, pmal-c, pFLAG, etc. The insertion into a cloning vector can, for example, be accomplished by ligating the DNA fragment into a cloning vector that has complementary cohesive termini. However, if the complementary restriction sites used to fragment the DNA are not present in the
15 cloning vector, the ends of the DNA molecules may be enzymatically modified. Alternatively, any restriction site desired may be produced by ligating nucleotide sequences (linkers) onto the DNA termini; these ligated linkers may comprise specific chemically synthesized oligonucleotides encoding restriction endonuclease recognition sequences. In addition, simple PCR or overlapping PCR may be used to
20 insert a fragment into a cloning vector.

Recombinant molecules can be introduced into host cells via transformation, transfection, infection, electroporation, etc., so that many copies of the gene sequence are generated. Preferably, the cloned gene is contained on a shuttle vector plasmid, which provides for propagation in a cloning cell, *e.g.*, *E. coli*, and
25 facile purification for subsequent insertion into an appropriate expression cell line, if such is desired. For example, a shuttle vector, which is a vector that can replicate in more than one type of organism, can be prepared for replication in both *E. coli* and *Saccharomyces cerevisiae* by linking sequences from an *E. coli* plasmid with sequences from the yeast 2 Φ plasmid.

30 In a preferred embodiment of the invention, the hNaIII18 sodium channel is cloned using a strategy designed to minimize mutations during cDNA

preparation, RT-PCR amplification, and growth in bacteria. This strategy is described in detail *infra*, in Example 1. The main points are summarized as follows:

First, as an alternative to conventional reverse transcriptases, which function optimally at temperatures of between 37 °C and 43 °C, this method employs
5 an avian RNase (-) reverse transcriptase that functions optimally at temperatures between 50-65 °C. The higher temperature serves to decrease secondary structure of the RNA to produce higher cDNA yield.

Second, for amplification of the cDNA, an enzyme mixture comprising the conventional thermostable Taq polymerase and Pwo polymerase is used. This
10 mixture is optimized to produce very large PCR products with low error frequency, thus decreasing the mutation frequency.

Third, the number of cycles of amplification is decreased to about 28, as opposed to the typical 30-35 cycles to further reduce the possibility of mutation.

Fourth, the PCR products are electrophoresed and visualized on an
15 agarose gel containing Crystal Violet stain, as opposed to ethidium bromide. Crystal Violet allows visualization in white light, eliminating the need for UV exposure. UV is known to induce mutations in ethidium bromide-stained DNA.

Fifth, to minimize recombination and mutation in plasmid DNA during amplification in bacteria, the PCR amplified cDNA is cloned into a low-copy number
20 expression vector that is engineered to have limited replication cycles and contains a tetracycline-resistance gene as a selectable marker instead of an ampicillin resistance gene. Fewer replication cycles again reduces the error rate during DNA synthesis, and selection with tetracycline is less likely to induce mutations in the plasmid than is ampicillin.

Sixth, competent bacterial cells that are designed to optimize cloning
25 of unstable inserts are selected for the transformation, and grown at a lower temperature (30-33 °C versus 37 °C) to decrease the growth rate and therefore, minimize the possibility of mutations. In addition, the cultures should be maintained in exponential (log) phase throughout growth, eliminating the possibility of mutations
30 resulting from starvation, poor aeration, and accumulation of toxic metabolites.

Seventh, small tetracycline resistant colonies are chosen for evaluation rather than large ones. Human NaIII expression during growth is expected to be toxic to bacteria, thus transformed cells will yield smaller colonies.

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hNaIII18 Regulatory Nucleic Acids

Elements of the hNaIII18 promoter can be identified by scanning the human genomic region upstream of the hNaIII18 start site, *e.g.*, by creating deletion mutants and checking for expression, or by using an algorithm. Sequences up to about 6 kilobases (kb) or more upstream from the hNaIII18 start site can contain
10 tissue-specific regulatory elements.

The term "hNaIII18 promoter" encompasses artificial or heterologous promoters. Such promoters can be prepared by deleting non-essential intervening sequences from the upstream region of the hNaIII18 promoter, or by joining upstream regulatory elements from the hNaIII18 promoter with a heterologous minimal
15 promoter, such as the CMV immediate early promoter.

A hNaIII18 promoter can be operably associated with a heterologous coding sequence, *e.g.*, for a reporter gene (luciferase and green fluorescent proteins are examples of reporter genes) in a construct. This construct can be used to test for conditions or reagents that normally result in expression. This construct can be used
20 in screening assays, described below, for hNaIII18 agonists and antagonists.

hNaIII18 regulatory nucleic acids of the present invention also include non-endogenous or artificial promoter sequences or sequences that encode zinc finger proteins that may be used, *e.g.*, in gene activation techniques, to initiate expression of hNaIII18 in cells where it is not normally expressed or to upregulate expression of the
25 hNaIII18 subunit protein to a higher level where it would otherwise be expressed in suboptimal levels. Gene activation techniques that may be adapted to this use are described in the art, *e.g.*, in U.S. Patent Nos. 5,968,502 and 6,214,622 to Treco et al.

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Expression of hNaIII18 Polypeptides

The primary goal for establishing a stable cell line expressing functional human sodium channels is to identify antagonists to inhibit sodium currents

mediated by the sodium channels. DRG neurons transmit nociceptive signals from the peripheral nervous system to the central nervous system. TTX-S and TTX-R sodium channels mediate the DRG action potentials responsible for these signals. However, DRG neurons express several different isoforms of TTX-S and TTX-R currents, thereby making it difficult to determine specific interactions of antagonists with particular subtypes of sodium channels in these cells.

By generating a cell line that expresses a single sodium channel subtype, *e.g.*, hNaIII18, alone or preferably in combination with appropriate β subunits, the effect of drugs on the different sodium channel isoforms can be assessed. Previously, developing stable cell lines expressing nucleic acids containing repetitive sequences, such as those contained within sodium channel genes, has been challenging. In particular, cell lines expressing functional sodium channels have been difficult to generate due to the occurrence of inactivating mutations arising in the cDNA during the cloning process (*i.e.*, cDNA preparation, PCR amplification, and subsequent growth in bacteria). International PCT publication WO 98/38302 (Delgado et al.) describes isolation, cloning and expression of a rat TTX-S sodium channel in *Xenopus* oocytes. Experiments described therein demonstrate the formation of a functional TTX-S channel after injection of cRNA into *Xenopus* oocytes for the α -subunit, alone or in combination with the β 1, β 2 or β 3 subunits. International PCT Publication WO 01/68681 (Aitken et al.) describes altered ion channel proteins having acquired sensitivity or refractory sensitivity to a gating agent. A rat sodium channel type II was modified by site-directed mutagenesis and PCR to contain sequences that bind α -scorpion toxins, which inactivate sodium channels, for use to evaluate ion channel activity and to screen for compounds for therapeutic applications. The modified sodium channel was then stably or transiently expressed in several mammalian host cells, including HEK293 variants and CHO cells, which were used in a high-throughput, plate-based screening assay.

International PCT publication WO/02068 (Korsgaard) describes stable cloning of a splice variant of a rat α I sodium channel in HEK293 cells.

To date, there have been no reports of stable expression of a cloned human sodium type III channel in mammalian cells. The method described herein combines several procedures to facilitate the cloning and generation of stable cell

lines containing such repetitive sequences, resulting in functional expression of such genes. In particular, the present invention describes the cloning and stable expression of a novel splice variant of human NaIII, designated hNaIII18.

5 The nucleotide sequence coding for hNaIII18, or an antigenic fragment, derivative or analog thereof, (including, *e.g.*, a chimeric protein) can be inserted into an appropriate expression vector, *i.e.*, a vector which contains the necessary elements for the transcription and translation of the inserted protein-coding sequence. Thus, a nucleic acid molecule having a nucleotide sequence encoding the hNaIII18 subunit protein of the invention can be operationally associated with a
10 promoter in an expression vector of the invention. Either a cDNA or genomic sequence can be cloned and expressed under control of such regulatory sequences. Such vectors can be used to express functional, or functionally inactivated, hNaIII18 polypeptides.

The necessary transcriptional and translational signals can be provided
15 on a recombinant expression vector, or they may be supplied from the native gene encoding hNaIII18 and/or its flanking regions.

Potential host-vector expression systems include but are not limited to mammalian cell systems transfected with expression plasmids or infected with virus (*e.g.*, vaccinia virus, adenovirus, adeno-associated virus, herpes virus, etc.); insect cell
20 systems infected with virus (*e.g.*, baculovirus); microorganisms such as yeast containing yeast vectors; and bacteria transformed with bacteriophage, DNA, plasmid DNA, or cosmid DNA. The expression elements of vectors vary in their strengths and specificities. Depending on the host-vector system utilized, any one of a number of suitable transcription and translation elements may be used.

25 Expression of the hNaIII18 protein may be controlled by any promoter/enhancer element known in the art, but these regulatory elements must be functional in the host selected for expression. Promoters which may be used to control hNaIII18 gene expression include, but are not limited to, cytomegalovirus (CMV) promoter (see, *e.g.*, U.S. Patent Nos. 5,385,839 and 5,168,062), the SV40
30 early promoter region (Benoist and Chambon, Nature 1981; 290:304-310), the promoter contained in the 3' long terminal repeat of Rous sarcoma virus (Yamamoto, *et al.*, Cell, 1980; 22:787-797), the herpes thymidine kinase promoter (Wagner *et al.*,

Proc. Natl. Acad. Sci. U.S.A., 1981; 78:1441-1445), the regulatory sequences of the metallothionein gene (Brinster *et al.*, Nature, 1982; 296:39-42, prokaryotic expression vectors such as the β -lactamase promoter (Villa-Komaroff, *et al.*, Proc. Natl. Acad. Sci. U.S.A. 1978; 75:3727-3731), or the tac promoter (DeBoer, *et al.*, Proc. Natl. Acad. Sci. U.S.A. 1983; 80:21-25) (see also "Useful proteins from recombinant bacteria" in Scientific American, 1980, 242:74-94), promoter elements from yeast or other fungi such as the Gal 4 promoter, the ADC (alcohol dehydrogenase) promoter, PGK (phosphoglycerol kinase) promoter, alkaline phosphatase promoter, and transcriptional control regions that exhibit tissue specificity, such as, *e.g.*, endothelial cell-specific promoters.

Solubilized forms of the protein can be obtained where necessary by solubilizing inclusion bodies or reconstituting membrane components, *e.g.*, by treatment with detergent, and if desired sonication or other mechanical processes, as described above. The solubilized protein can be isolated using various techniques, such as polyacrylamide gel electrophoresis (PAGE), isoelectric focusing, 2-dimensional gel electrophoresis, chromatography (*e.g.*, ion exchange, affinity, immunoaffinity, and sizing column chromatography), centrifugation, differential solubility, immunoprecipitation, by any other standard technique for the purification of proteins, or by a combination of such techniques.

Since β -subunits 1-3 are known to bind the α -subunits of sodium channels, the present invention also contemplates co-expression of a β -subunit with NaIII18. While the role played by β -subunits in determining the pharmacological properties of voltage-gated sodium channels appears to be minor, at least for the commonly-studied binding sites, the β -subunits do appear to have effects on the biophysics (gating kinetics) of sodium channel function. Therefore, to the extent that biophysics and drug interactions are linked, the β -subunits may affect pharmacology of agents used to modulate sodium channel activity. Some known β -subunits that may be co-expressed with the NaIII18 subunit of the invention are described in Isom *et al.*, Neuron 1994; 12:1183-94; International PCT publication WO 01/44293 to Plumpton *et al.*; International PCT publication WO 01/23570 to d'Andrea *et al.*; U.S. published patent application 2002/0045229 to Qin *et al.*; and under GenBank Accession Nos.

U87445, AF007783, AH005825, AF007783, AF04948, L10338 and L16242, among others

hNaIII18 Binding Partners

5 The present invention further provides a method for identifying physiological binding partners of hNaIII18. One method for evaluating and identifying hNaIII18 binding partners is the yeast two-hybrid screen. Preferably, the yeast two-hybrid screen is performed using a cell library with yeast that are transformed with recombinant hNaIII18. Alternatively, hNaIII18 can be used as a
10 capture or affinity purification reagent. In another alternative, labeled hNaIII18 can be used as a probe for binding, *e.g.*, by immunoprecipitation or Western analysis. Several expected hNaIII18 binding partners are the sodium channel β subunits, as described in the section above.

 Generally, binding interactions between hNaIII18 and any of its
15 binding partners will be strongest under conditions approximating those found in the native cell, *i.e.*, physiological conditions of ionic strength, pH and temperature, and particularly those obtaining in the cell membrane. Perturbation of these conditions will tend to disrupt the stability of a binding interaction.

Antibodies to hNaIII18

 Antibodies to hNaIII18 are useful, *inter alia*, for determining the presence of hNaIII18 in a cell and for cellular regulation (*i.e.*, inhibition) of hNaIII18 activity, as set forth below. According to the invention, a hNaIII18 polypeptide produced recombinantly or by chemical synthesis, and fragments or other derivatives
25 or analogs thereof, including fusion proteins, may be used as immunogens to generate antibodies that recognize the hNaIII18 polypeptide. Such antibodies include but are not limited to polyclonal, monoclonal, chimeric, single chain, Fab fragments, and Fab expression libraries. Such an antibody binds specifically to hNaIII18, and may recognize either a mutant form of hNaIII18 or wild-type hNaIII18, or both. The
30 antibodies of the present invention are specific for hNaIII18 and either do not recognize, or bind with lower affinity to, orthologs of hNaIII18. In one embodiment,

specific binding of such antibodies to hNaIII18 polypeptides provides the ability to detect the presence of the hNaIII18 polypeptide in a sample. In another embodiment, specific binding of such antibodies to hNaIII18 polypeptides provides the ability to preferentially inhibit the activity of hNaIII18, or an ion channel comprising hNaIII18.

5 Various procedures known in the art may be used for the production of antibodies against hNaIII18 polypeptides. These include but are not limited to the hybridoma technique originally developed by Kohler and Milstein (Nature 1975; 256:495-497), as well as the trioma technique, the human B-cell hybridoma technique (Kozbor *et al.*, Immunology Today 1983, 4:72; Cote *et al.*, Proc. Natl. Acad. Sci. 10 1983, 80:2026-2030), and the EBV-hybridoma technique to produce human monoclonal antibodies (Cole *et al.*, in Monoclonal Antibodies and Cancer Therapy, Alan R. Liss, Inc., 1985, pp. 77-96).

hNaIII18 Agonists and Antagonists

15 The present invention also contemplates the identification of compounds that modulate hNaIII18 sodium channel activation and activity. Such compounds are useful, *e.g.*, for inhibiting (*i.e.*, antagonizing) or increasing (*i.e.*, agonizing) biological activities that are associated with sodium channel activation and/or as therapeutic agents for treating disorders associated with excessive sodium channel activation.

20 Compounds that modulate hNaIII18 activity or an activity associated therewith may be readily identified using screening methods of the present invention. In one embodiment, compounds identified by the screening methods of this invention bind to a hNaIII18-subunit containing ion channel. Compounds identified by the present method may antagonize or agonize hNaIII18 subunit-containing channel 25 activity, as well as a related downstream biological effect (*e.g.*, the ability of DRG to transmit nociceptive signals from the PNS to the CNS) that are associated with excessive sodium channel current and activity.

30 *In vivo* or cell culture assays may be used to determine whether a test compound functions as an antagonist to inhibit hNaIII18 activity in cells. For instance, cell culture assays may be used to measure a test compound's ability to modulate an activity, such as induction, strength or duration of sodium channel

current associated with hNaIII18 subunit-containing sodium channel activity. Such assays generally comprise contacting a cell that expresses a hNaIII18 subunit containing sodium channel with a test compound. The cell should preferably be contacted with the test compound before or during exposure to an agent or stimulus that otherwise would serve to depolarize the cell membrane and thus activate (*i.e.*, open) the sodium channel: *e.g.* a high potassium chloride saline solution, or an extracellular field-stimulating electrode. The cell can then be examined to determine whether a response otherwise associated with sodium channel activation has been inhibited. In a non-limiting embodiment, the response of the cell treated with the test compound is compared to that of a control cell that has not been treated with the test compound. Cell assays include those utilizing conventional, electrode-based, electrophysiological techniques, as well as the new generation high-throughput, planar electrode (orifice) -based, electrophysiological technologies, among others. Other assays include monitoring changes in membrane potential with appropriate fluorescent, or luminescent, dyes, measuring ion flux through the sodium channel with a radiolabeled tracer, or assaying downstream consequences of sodium channel activation, such as calcium mobilization or effects on gene expression, using an appropriate reporter system.

Positive modulation (*i.e.*, agonism) of hNaIII18 subunit-containing channels may be desirable under certain circumstances, and screening for such agonists can be conducted according to the methods of the invention.

Screening

According to the present invention, nucleotide sequences encoding hNaIII18 are useful targets to identify drugs that are effective in preventing or alleviating pain, or drugs that can be used as anti-epileptics/anticonvulsants, anesthetic antiarrhythmics, and in the treatment of bipolar disorder (see section entitled Therapeutics, below), any of which may be associated with the function of the sodium channel. Examples of such drugs include without limitation: (i) isolated nucleic acids capable of altering expression of hNaIII18 (*e.g.*, antisense or ribozyme molecules); (ii) small organic molecules that bind to and modulate the function of a hNaIII18 subunit or a hNaIII18 subunit-containing ion channel; and (iii) peptides or

peptide analogs that bind to and modulate the function of a hNaIII18 subunit or a hNaIII18 subunit-containing ion channel. In addition, the nucleotide sequences encoding hNaIII18 are useful for studying the role of the channels both in pain perception and in physiological and pathological brain functions.

5 Any screening technique known in the art can be used to screen for agonists or antagonists. The present invention contemplates screens for small molecules and mimics, as well as screens for natural products that bind to and agonize or antagonize hNaIII18-containing ion channels. For example, natural product libraries can be screened using assays of the invention for molecules that agonize or
10 antagonize hNaIII18-containing ion channel activity.

Knowledge of the primary sequence of hNaIII18, and the similarity of that sequence with proteins of known function, can provide an initial lead to inhibitors or antagonists. Identification and screening of modulators is further facilitated by determining structural features of the protein, *e.g.*, using X-ray crystallography,
15 neutron diffraction, nuclear magnetic resonance spectrometry, and other techniques for structure determination. These techniques provide for the rational design or identification of agonists and antagonists.

Another approach uses recombinant bacteriophage to produce large libraries. Using the "phage method" (Scott and Smith, Science 1990, 249:386-390; Cwirla, et al., Proc. Natl. Acad. Sci. USA 1990, 87:6378-6382; Devlin et al., Science
20 1990, 49:404-406), very large libraries can be constructed (10⁶-10⁸ chemical entities). A second approach uses primarily chemical methods, of which the Geysen method (Geysen et al., Molecular Immunology 1986, 23:709-715; Geysen et al. J. Immunologic Methods 1987, 102:259-274); and the method of Fodor et al. (Science
25 1991, 251:767-773) are examples. Furka et al. (14th International Congress of Biochemistry 1988, Volume #5, Abstract FR:013; Furka, Int. J. Peptide Protein Res. 1991, 37:487-493), Houghton (U.S. Patent No. 4,631,211) and Rutter et al. (U.S. Patent No. 5,010,175) generally describe methods to produce a mixture of peptides that can be tested as agonists or antagonists.

30 In another aspect, synthetic libraries, such as those described in Needels et al., Proc. Natl. Acad. Sci. USA 1993, 90:10700-4; Ohlmeyer et al., Proc. Natl. Acad. Sci. USA 1993, 90:10922-10926; Lam et al., PCT Publication No. WO

92/00252; and Kocis et al., PCT Publication No. WO 9428028, and the like, can be adapted to screen for compounds according to the present invention.

Test compounds can be screened from large libraries of synthetic or natural compounds. Numerous means are currently used for random and directed
5 synthesis of saccharide, peptide, and nucleic acid based compounds. Synthetic compound libraries are commercially available from a variety of sources, including Maybridge Chemical Co. (Trevillet, Cornwall, UK), Comgenex (Princeton, NJ), Brandon Associates (Merrimack, NH), and Microsource (New Milford, CT). A rare chemical library is available from Aldrich (Milwaukee, WI). Alternatively, libraries
10 of natural compounds in the form of bacterial, fungal, plant and animal extracts are available from a variety of sources including, *e.g.*, Pan Laboratories (Bothell, WA) and MycoSearch (NC), or are readily producible *de novo*. Additionally, natural and synthetically produced libraries and compounds are readily modified through conventional chemical, physical, and biochemical means (see, *e.g.*, Blondelle et al.,
15 TIBTech 1996, 14:60).

In Vitro Screening Methods and Activity Assays

Cell-based screening

Intact cells expressing a hNaIII18 subunit-containing ion channel can
20 be used in screening methods to identify candidate compounds useful in modulating the activity of sodium channels containing hNaIII18. In one embodiment, a cell line is established that stably expresses or overexpresses the hNaIII18 subunit protein, either alone or in combination with one or more other sodium channel β subunits, to form a functional sodium channel. Alternatively, cells (including without limitation
25 mammalian, invertebrate, yeast, or bacterial cells) are transiently programmed to express a hNaIII18 subunit protein by introduction of the appropriate DNA or mRNA. Identification of candidate compounds can be achieved using any suitable assay, including without limitation: (i) assays that measure binding of test compounds to hNaIII18 (alone or in combination with sodium channel β subunits described *supra*):
30 (ii) assays that measure the ability of a test compound to modulate (*i.e.*, agonize or antagonize) a measurable activity or function of hNaIII18 or a hNaIII18 subunit-containing ion channel; and (iii) assays that measure the ability of a compound to

enhance or inhibit the transcriptional activity of sequences derived from the promoter (*i.e.*, regulatory) regions of the hNaIII18 gene.

Any cell assay system that allows for assessment of functional activity of a hNaIII18 subunit-containing sodium channel is encompassed by the present invention. In a specific embodiment, described *infra*, the assay can be used to identify compounds that selectively modulate the hNaIII18 subunit protein, which can be determined by assessing the effects on NaIII18 subunit-expressing cells contacted with a test compound. The assay system can thus be used to identify compounds that selectively produce a functional effect through hNaIII18 sodium channels.

Compounds that decrease activity of the sodium channel in response to activation may be useful as novel therapeutics in the amelioration of neuropathic pain mediated by DRG neurons, or as anti-epileptics/convulsants, anesthetics, antiarrhythmics, or in the treatment of bipolar disorder.

Compounds that increase activity of sodium channels may be useful as cognitive enhancers, or in disorders such schizophrenia. In these instances, a subtype-selective agent would be preferable to offset the potential for proconvulsant effects and to increase cardiac contractility in individuals suffering from heart failure.

Alternatively, the change in membrane potential induced by sodium ions of the voltage-gated channel-containing cells may be monitored using fluorescence methods. When using fluorescence methods, the voltage-gated channel containing cells may be incubated with a membrane potential indicating agent that allows for a determination of changes in the membrane potential of the cells caused by the influx of sodium ions. Such membrane potential indicating agents include fluorescent indicators, such as those provided in a Molecular Devices Membrane Potential Kits for the FLIPR/Flexstation, DIBAC4(3), DiOC6(6) DiOC5(3), DiOC2(3) and fluorescence resonance energy transfer (FRET) based dyes such as JC1, and JC9, among others.

Another method that allows for assessment of functional activity of hNaIII18-containing sodium channels involves monitoring the change in membrane potential induced by sodium ions on the channel-containing cells by fluorescent methods, *e.g.*, using a FLIPR assay (Fluorescence Image Plate Reader; available from Molecular Devices)(Rose et al. Pflugers Arch. 1999 Dec;439(1-2):201-7). Another

method involves radioactive flux assays that measure the ability of radioactive tracer ions such as [^{22}Na] and [^{14}C] guanidinium to pass into the cell upon channel activation (Barann M. et al. Naunyn Schmiedebergs Arch Pharmacol. 1999; 360(3):234-41).

After the channel is activated, concentrations of these tracer ions increase inside the cell. Free extra-cellular tracer is washed away, cells are lysed, and radioactivity in the lysates is counted using standard scintillation counters or other radioactivity analysis instruments.

Yet another method involves measuring cell viability upon veratridine-mediated stabilization of sodium channels in their open conformation (Okuyama K. et al., Eur J Pharmacol. 2000; 398(2):209-16). Cells undergo toxic sodium overload followed by cell death. Compounds that prevent cell death, or cellular toxicity, can be assayed with standard cytotoxicity kits and with standard cell viability dyes such as alamar blue.

Cell-Free Screening

In another embodiment, an assay is a cell-free assay comprising contacting a hNaIII18 polypeptide or biologically active portion thereof with a test compound and determining the ability of the test compound to modulate (*e.g.*, stimulate or inhibit) the activity of the hNaIII18 polypeptide or biologically active portion thereof.

In yet another embodiment, the cell-free assay comprises (i) contacting the hNaIII18 polypeptide of the invention or biologically active portion thereof with a known compound or polypeptide which binds the hNaIII18 polypeptide to form an assay complex; (ii) contacting the assay complex with a test compound; (iii) determining the ability of the test compound to interact with the hNaIII18 polypeptide by determining the ability of the test compound to modulate the effect of the known compound on the activity of the sodium channel.

More specifically, a cell-free method can involve monitoring the specific binding of a radiolabeled sodium channel selective neurotoxin, such as [^3H]tetrodotoxin or [^3H]batrachotoxin, or a high affinity small-molecule ligand, to a membrane preparation from cells or tissues engineered to express hNaIII18-containing sodium channels (Garritsen A. et al. Eur J Pharmacol. 1988; 145(3):261-6;

MacKinnon AC. et al. J Pharmacol. 1995; 115(6):1103-9; Bambrick L. et al., J Pharmacol Toxicol Methods. 1994; 32(3):129-38). Following techniques that are well know in the art, total binding to membranes can be measured upon incubation with the radioligand until the biomolecular reaction reaches equilibrium. Nonspecific binding is defined in the presence of an unlabelled competitor ligand. Specific binding is the subtraction of total minus nonspecific binding. Compounds that modulate specific binding can thereby be identified.

In another embodiment, modulators of expression of the hNaIII18 polypeptide of the invention are identified in a method in which a cell is contacted with a candidate compound and the expression of the mRNA or protein corresponding to hNaIII18 in the cell is determined. The level of expression of the hNaIII18 mRNA or protein in the presence of the candidate compound is compared to the level of expression of the hNaIII18 mRNA or protein in the absence of the candidate compound. The candidate compound can thereby be identified as a modulator of expression of the hNaIII18 polypeptide of the invention based on this comparison. For example, when expression of the hNaIII18 mRNA or protein is increased in the presence of the candidate compound compared to in the absence of the candidate compound, then the candidate compound is identified as a stimulator of hNaIII18 mRNA or protein expression. Alternatively, when expression of the hNaIII18 mRNA or protein is specifically reduced in the presence of the candidate compound compared to in the absence of the candidate compound, then the candidate compound is identified as an inhibitor of hNaIII18 mRNA or protein expression. In view of this disclosure, the level of the hNaIII18 mRNA or protein expression in cells can be determined by methods known in the art.

High-Throughput Screen

Drug candidates according to the invention can be identified by screening in high-throughput assays, including without limitation cell-based or cell-free assays. It will be appreciated by those skilled in the art that different types of assays can be used to detect different types of drug candidates. Several methods of automated assays have been developed in recent years so as to permit screening of tens of thousands of compounds in a short period of time. Such high-throughput

screening methods are particularly preferred. The use of high-throughput screening assays to test for agents is greatly facilitated by the availability of the large amounts of purified hNaIII18 polypeptides provided by the invention.

5

Therapeutic Uses

It is desirable to modulate the function of sodium channels in a number of clinical and therapeutic environments. Sodium channels are implicated in conditions including chronic and neuropathic pain, cardiac arrhythmias (Duch et al., Toxicol Lett 1998; 100-101:255-63), neuronal disorders associated with deficient
10 oxygen supply or mitochondrial dysfunction (Urenjak et al., Amino Acids 1998;14(1-3):151-8), and epilepsy (Ragsdale et al., Brain Res Rev 1998;26(1):16-28). In addition, inhibition of sodium channels is an effect of local anesthetics (Li et al., Mol Pharmacol 1999; 55(1):134-41).

According to the present invention, inhibition of hNaIII18 subunit-
15 containing sodium channel activity may be used as a treatment option in patients with a pain disorder, such as but not limited to a neuropathic pain-related disease such as, *e.g.*, pain from peripheral nerve trauma, herpes virus infection, diabetes mellitus, causalgia, plexus avulsion, neuroma, limb amputation, and vasculitis. Neuropathic pain is also caused by nerve damage from chronic alcoholism, human
20 immunodeficiency virus infection, hypothyroidism, uremia, or vitamin deficiencies. The neuronal hyperexcitability and corresponding molecular changes in neuropathic pain have many features in common with the cellular changes in certain forms of epilepsy. This has led to the use of anticonvulsant drugs for the treatment of neuropathic pain (Jensen, Eur J Pain 2002;6 Suppl A:61-8). Local anesthetics such as
25 lidocaine and mexiletine have also be shown to inhibit TTX-S sodium channel activity in hyperexcitable neurons in rat (Novartis Found Symp 2002;241:189-201; discussion 202-5, 226-32).

Inhibition of the sodium channel of the present invention may also be used as a treatment option in patients with chronic pain. In chronic pain, the pain can
30 be mediated by multiple mechanisms. This type of pain generally arises from injury to the peripheral or central nervous tissue. The chronic pain-type syndromes include pain associated with spinal cord injury, multiple sclerosis, post-herpetic neuralgia,

trigeminal neuralgia, phantom pain, causalgia, and reflex sympathetic dystrophy and lower back pain.

Inhibition of the sodium channel of the present invention may also be used as a treatment option in patients with nociceptive pain.

5

Inhibition of Protein Synthesis or Sodium Channel Activity

Gene transcription and protein translation may be inhibited by administration of exogenous compounds. Exogenous compounds may interact with extracellular and/or intracellular messenger systems to regulate protein synthesis. In this embodiment, exogenous compounds that inhibit hNaIII18 protein synthesis may be used in the prevention and/or treatment for pain resulting from persistent channel activity.

Accordingly, in an exemplary embodiment, the modulatory method of the invention involves contacting a cell, tissue or subject with an agent that modulates one or more of the activities of hNaIII18 protein activity associated with the cell. An agent that modulates hNaIII18 protein activity can be an agent as described herein, such as a nucleic acid or a protein, an hNaIII18-specific antibody, an hNaIII18 agonist or antagonist, a peptidomimetic of an hNaIII18 agonist or antagonist, or other small molecule. In one embodiment, the agent stimulates one or more hNaIII18 activities. In another embodiment the agent inhibits one or more hNaIII18 activities. Examples of such inhibitory agents include antisense hNaIII18 nucleic acid molecules, anti-hNaIII18 antibodies, and hNaIII18 inhibitors. These modulatory methods can be performed *in vitro* (e.g., by culturing the cell with the agent) or, alternatively, *in vivo* (e.g., by administering the agent to a subject). As such, the present invention provides methods of treating an individual afflicted with a disease or disorder characterized by aberrant or unwanted expression or activity of a hNaIII18 protein or nucleic acid molecule. In one embodiment, the method involves administering an agent (e.g., an agent identified by a screening assay described herein), or combination of agents that downregulates hNaIII18 expression or activity or the activity of a hNaIII18 subunit-containing ion channel.

30

In yet another embodiment, the agent enhances one or more hNaIII18 activities, such as by administering a hNaIII18 protein or nucleic acid molecule as therapy to compensate for reduced or aberrant hNaIII18 expression or activity.

5 The present invention further provides antisense nucleic acids, which may be used to inhibit expression of hNaIII18 nucleotide sequences of the invention. This antisense technology has been described as inhibiting the peripheral tetrodotoxin (TTX)-resistant sodium channel, NaV1.8, found in sensory neurons, when administered intrathecally (Lai et al., Pain 2002; 95 (1-2):143-52). According to this method, the antisense nucleic acid, upon hybridizing under cytoplasmic conditions
10 with complementary bases in an RNA or DNA molecule, inhibits the RNA or DNA. Additionally, hybridization of the antisense nucleic acid to the DNA or RNA may inhibit transcription of the DNA into RNA and/or translation of the RNA into the protein. If the RNA is a messenger RNA transcript, the antisense nucleic acid is a counter-transcript or mRNA-interfering complementary nucleic acid. Antisense
15 nucleic acid molecules can be encoded by a recombinant gene for expression in a cell (see, e.g., U.S. Patent No. 5,814,500; U.S. Patent No. 5,811,234) or can be prepared synthetically (e.g., U.S. Patent No. 5,780,607).

Alternatively, antibody molecules or antigen-binding antibody fragments can be administered either directly or by expressing nucleotide sequences
20 encoding antibodies or binding fragments thereof within the target cell population by utilizing, for example, techniques such as those described in Marasco *et al.* (Proc. Natl. Acad. Sci. USA, 1993, 90:7889-7893).

Formulations and Administration

25 The drug candidate or agent that modulates hNaIII18 activity is advantageously formulated in a pharmaceutical composition by admixing the drug candidate or agent with a pharmaceutically acceptable carrier. This agent may then be designated as the active ingredient, or therapeutic agent for use, for example, against chronic, neuropathic pain, or nociceptive pain

30 The form, amount and route of administration of the therapeutic compound envisioned for use depends on the type and severity of the disease or condition to be treated, as well as the patient's state of health, gender, weight, age,

etc., and can be determined by an attending medical practitioner in view, *e.g.*, of the results of published clinical trials. The concentration or amount of the active ingredient depends on the desired dosage and administration regimen, as discussed below. Suitable dose ranges may include from about 1 mg/kg to about 100 mg/kg of body weight per day.

The pharmaceutical compositions may also include other biologically active substances in combination with the NaIII18 modulatory agent. Such substances include but are not limited to opioids such as morphine, codeine, fentanyl, oxycodone, hydrocodone, and buprenorphine; and non-steroidal anti-inflammatory drugs (NSAID's) such as but not limited to ibuprofen and COX-2 inhibitors, among others

The phrase "pharmaceutically acceptable" refers to molecular entities and compositions that are physiologically tolerable and do not typically produce an allergic or similar untoward reaction, such as gastric upset, dizziness and the like, when administered to a human. Preferably, as used herein, the term "pharmaceutically acceptable" means that the carrier has been approved by a regulatory agency of the Federal or a state government or listed in the U.S. Pharmacopeia or other generally recognized pharmacopeia for use in animals, and more particularly in humans. The term "carrier" refers to a diluent, adjuvant, excipient, or vehicle with which the active ingredient is administered. Such pharmaceutical carriers can be sterile liquids, such as water and oils, including those of petroleum, animal, vegetable or synthetic origin, such as peanut oil, soybean oil, mineral oil, sesame oil and the like. Water or aqueous solution saline solutions and aqueous dextrose and glycerol solutions are preferably employed as carriers, particularly for injectable solutions. Suitable pharmaceutical carriers are described in "Remington's Pharmaceutical Sciences" by E.W. Martin.

According to the invention, the pharmaceutical composition of the invention can be introduced parenterally, transmucosally, *e.g.*, orally (per os), nasally, rectally, or transdermally. Parental routes include intravenous, intra-arteriole, intramuscular, intradermal, subcutaneous, intraperitoneal, intraventricular, and intracranial administration. The pharmaceutical composition may alternatively be

adapted for topical or transdermal application, such in a salve, cream, lotion, spray or transdermal patch system.

The pharmaceutical compositions may be added to a retained physiological fluid such as blood or synovial fluid. For CNS (Central Nervous System) administration, a variety of techniques are available for promoting transfer of the therapeutic across the blood brain barrier including disruption by surgery or injection, co-administration of drugs that transiently open adhesion contact between CNS vasculature endothelial cells, and co-administration of substances that facilitate translocation through such cells.

In another embodiment, the active ingredient can be delivered in a vesicle, in particular a liposome (see Langer, Science 1990; 249:1527-1533; Treat et al., in Liposomes in the Therapy of Infectious Disease and Cancer, Lopez-Berestein and Fidler (eds.), Liss: New York 1989 pp. 353-365; Lopez-Berestein, *ibid.*, pp. 317-327; see generally *ibid.*).

In yet another embodiment, the therapeutic substance can be delivered in a controlled release formulation. For example, an active ingredient may be administered using intravenous infusion with a continuous pump, in a polymer matrix such as poly-lactic/glutamic acid (PLGA), a pellet containing a mixture of cholesterol and the active ingredient (Silastic^{RTM}; Dow Corning, Midland, MI; see U.S. Patent No. 5,554,601) implanted subcutaneously, an implantable osmotic pump, a transdermal patch, liposomes, or other modes of administration.

Compounds identified in the screening methods described herein (*i.e.*, modulators of sodium channel activity), may be provided to the patient in formulations that are known in the art and may include any pharmaceutically acceptable additives, such as excipients, lubricants, diluents, flavorants, colorants, and disintegrants. The formulations may be produced in useful dosage units such as tablet, caplet, capsule, liquid, or injection. In a further embodiment, these compounds are also administered in conjunction with other therapeutic agents such as the local anesthetics and anti-epileptic or anti-convulsants discussed *supra*.

The form and amount of therapeutic compound envisioned for use depends on the type of disease and the severity of the desired effect, patient state, etc., and can be determined by one skilled in the art.

EXAMPLES

The present invention is also described by means of an example, presented below. The use of such an example is illustrative only and in no way limits the scope and meaning of the invention or of any exemplified term. Likewise, the invention is not limited to any particular preferred embodiments described herein. Indeed, many modifications and variations of the invention will be apparent to those skilled in the art upon reading this specification and can be made without departing from its spirit and scope. The invention is therefore encompassed by the appended claims along with the full scope of equivalents to which the claims are entitled.

EXAMPLE 1: CLONING AND EXPRESSION OF HUMAN NaIII18

Methods

Reverse transcription and amplification of hNaIII18 cDNA. Reverse transcription was carried out using ThermoScript Reverse Transcriptase (Life Technologies, Rockville, MD), at an annealing temperature of 55 °C to maximize the likelihood of obtaining a full-length mRNA, according to manufacturer's instructions.

The following primers were designed to amplify the resulting full-length hNaIII18 cDNA:

forward primer (SEQ ID NO: 7)	5' - ATAAGAATGCGCCGCTGAAAAGATGGCACAGGCAC-3'
reverse primer (SEQ ID NO: 8)	5' - ATAGTTTAGCGGCCGCCTTGAAGTCCAGTTGACACA -3'

Primers were designed from the human NaIII (SCN3A) mRNA sequence previously identified (GenBank Accession # AJ251507).

Full-length cDNA (6000 base-pairs) was amplified using the Expand Long Template PCR (Boehringer Mannheim, Indianapolis, IA) according to the manufacturer's instructions. This enzyme is a mixture of thermostable Taq and Pwo

DNA polymerases. The number of cycles used for amplification was decreased to 28 cycles instead of the traditional 30-35 as an added precaution to minimize the occurrence of mutations during PCR.

Purification and cloning of PCR products into expression vectors.

5 PCR products resulting from the above-described reaction were visualized after electrophoresis on an agarose gel containing Crystal Violet. DNA was purified from the gel using methods well known in the art. DNA was stored in Tris-EDTA buffer, pH 7.4.

10 The PCR-amplified cDNA was cloned into a low-copy number expression vector, pLCTM1 (kindly provided by Al Goldin, UCI) according to standard procedures. This vector is under the control of the origin of replication (ORI) from plasmid pACYC184, which has a limited number of replication cycles, resulting in a decreased error rate during DNA replication.

15 Further, the plasmid contains a tetracycline-resistance gene instead of an ampicillin-resistance gene for selection. Tetracycline is less likely to induce mutations than ampicillin during selection. The plasmid also contains a neomycin resistant gene (NeoR) for selection of stable cell lines using the neomycin analog G418.

20 Once cloned, the vectors were transformed into maximum efficiency STBL2 competent *E. coli* bacteria (Life Technologies, Rockville, MD), provided in the kit according to manufacturer's instructions. These cells optimize the cloning of unstable inserts. Bacteria expressing hNaIII18 were grown at 30-33°C, and maintained in exponential (log) growth phase for the duration of culture.

25 Small tetracycline-resistant colonies were selected and grown-up for small-scale DNA preparations and large-scale preparations. The concentration of tetracycline was kept low (15 µg/ml) to further minimize adverse growth conditions. The cDNA was extracted using the Wizard Plus SV Minipreps DNA Purification System Kit (Promega, Madison, WI) according to the manufacturer's instructions, or Qiagen Midipreps according to manufacturer's instructions (Qiagen, Valencia, CA).
30 cDNA was then analyzed by restriction digest, and partial sequencing. Full sequencing was performed by MWG (North Carolina). Partial sequencing was done with standard DTCS sequencing method using a commercial Beckman Coulter kit.

Transient and stable transfection. In order to identify functional clones, human embryonic kidney cells (HEK293) were transiently transfected with clones that were identified as having the correct insert, and surveyed by an electrophysiological assay (Fugene transfection reagent, according to manufacturer's recommendation). One clone, pLCTM1huNaIII-18, was determined to be functional as it gave large TTX-S currents with the expected activation and inactivation kinetics typical of NaIII channel. For example, typical activation is measured within fractions of ms at $V_m=0\text{mV}$ (corresponding I_{max}). Inactivation is measured as the time constant between 1-3 ms at $V_m=0\text{mV}$ (increasing to 20 ms at -50mV to 0.5 ms at $+40\text{mV}$). Recovery from inactivation is a time constant of about 10ms at $V_m=-100\text{mV}$ and 60 ms at -80mV (see *e.g.*, Cummins et al., J Neurosci 2001; 21:52-5961).

This clone was fully sequenced for confirmation. In addition, several non-functional clones were partially sequenced.

Clone pLLCTM1huNaIII-18 was used to generate a stable cell line in HEK293 cells. Fugene-mediated transfection of HEK cells was performed in 35 mm dish followed by G418 selection (300 and 500 $\mu\text{g/ml}$), colony isolation, line expansion. G418-resistant cells were then analyzed with immunocytochemistry, RT-PCR and electrophysiology according to standard techniques.

Electrophysiology. Stably transfected cells were grown on poly DL-lysine-coated glass coverslips at $\sim 2,000$ cells/slip, or Petri dishes at $\sim 10,000$ cells/dish and were then placed into the electrophysiology recording chamber and infused with an extracellular solution (140 mM NaCl, 4.7 mM KCl, 1.2 mM MgCl_2 , 1 mM CaCl_2 , 11 mM glucose and 5 mM HEPES, pH 7.4) at a rate of 2 ml/min. Electrodes were prepared by pulling Patch pipettes (borosilicate glass) using a Sutter P-97 electrode puller, and were filled with a solution containing 110 mM CsCl, 10 mM NaCl, 5 mM MgCl_2 , 11 mM EGTA, 10 mM HEPES, 2 mM ATP and 1 mM GTP, pH 7.25, osmolarity 275-290 mOsm. When filled with this solution, the electrodes had resistances of about 1-4 MS. Currents were recorded using a whole-cell voltage clamp techniques as described in Hamill et al. (Pflugers Arch. 1981; 391; 85-100), at room temperature (21-23 °C). Briefly, currents were recorded using an Axopatch 200A amplifier (Axon Instruments, Foster City, CA) and were leak-subtracted (P/4),

low-pass filtered (3 kHz, 8-pole Bessel), digitized (20-50- μ s intervals), and stored using Digidata 1200 B interface and Pclamp6/Clampex software (Axon Instruments, Foster City, CA). Residual series access resistance was largely (75-80%) canceled using built-in amplifier circuitry. The junction potential calculated using JPCalcW software (Cell MicroControl, Virginia Beach, VA) was small (<7 mV); so, no correction of the holding voltage was made.

To take I-V curves, cells were held at a holding voltage, $V_h = -90$ mV. A series of 16 depolarizing pulses (10 ms in duration) incrementing in 10 mV steps were applied at a frequency of 0.5 Hz. The peak values of currents were plotted against corresponding voltage steps to get the I-V curve. From this plot V_{max} , *i.e.*, the voltage causing the maximal Na^+ current, as well as rising times to peak and time constant for inactivation at different voltages were determined. To get steady-state inactivation curves, cells were held at a holding voltage, $V_h = -120$ mV to remove residual inactivation. A series of 30 depolarizing conditioning pre-pulses (each 100 ms in duration) incrementing in 5 mV steps immediately followed by a 5 ms testing pulse, V_t , to V_{max} were applied at a frequency of 0.5 Hz. The peak currents in response to V_t were plotted against the size of corresponding conditioning pre-pulses, V_c , to get steady-state inactivation curve. The Boltzman fit to this curve, *i.e.*, $\{1/[1+\exp((V+V_{1/2})/k)]\}$, returned the values of $V_{1/2}$ (the half-inactivation voltage) and k (the slope of the curve).

To measure recovery from inactivation, cells were held at a holding voltage $V_h = -120$ mV to remove residual steady-state inactivation. The depolarizing conditioning pre-pulse (100 ms in duration) was applied to V_c to cause complete inactivation of the channels (usually $V_c = -10$ mV). The conditioning pre-pulse was immediately followed by hyperpolarizing gap back to -120 mV of a variable duration. The gap duration was incremented in subsequent cycles in varying steps (2 ms -100 ms) depending on the speed of recovery. The gap was immediately followed by the testing pulse V_t (10 ms in length) to assess the fraction of Na^+ channels available for activation. The cycle was repeated every 5 seconds while the gap duration was incremented. The peak currents to V_t were plotted against the corresponding gap

duration to get the kinetics of recovery. The mono- or double- exponential fit to the data returned the time constant, $\tau_{\text{repr.}}$, of repriming from inactivation.

Results

5 **Identification of a splice-variant for human NaIII (SCN3).** Clone pLCM1huNaIII-18 is a novel splice variant and contains an additional 147 nucleotides corresponding to 49 amino acids in the cytoplasmic loop between domain 1S6 and IIS1 (see SEQ ID NO: 1 and SEQ ID NO: 2). Partial sequencing of several other clones that were not determined to have functional activity revealed sequences
10 that either matched the published sequence (GenBank Accession #AJ251507) or contained an extra 9 or 96 nucleotides. The shorter splicing patterns correspond to what had been described for the rat NaIII clone (Schaller et al., *J Neurosci* 1992; 12(4):1370-81), resulting in a protein with an additional 3 (rNaIIIa) or 22 (rNaIIIb) amino acids, but had not been described for the human NaIII before.

15 Subsequent to the completion of the cloning of hNaIII18, it was discovered that a clone having the same 147 nucleotide insert was deposited in GenBank on February 1, 2001 (GenBank Accession # AF225986-SEQ ID NO: 5). See cDNA alignment in Figure 8. However, that encoded amino acid sequence differs from the sequence disclosed herein by 12 amino acids (between two clones), at
20 amino acid residues 208, 475, 495, 508, 604, 1163, 1576, 1614, 1741, 1743, 1862 and 1966, respectively (SEQ ID NO: 2 vs. SEQ ID NO: 6). See amino acid alignment of Figure 9.

 Stable transfection of the pLCM1huNaIII-18 resulted in the generation of two cell lines that expressed the expected ~220 kDa hNaIII18 protein and
25 exhibited functional sodium channels, designated 293/huNaIII18-300-20 and 293/huNaIII18-500-35, with appropriate TTX-S currents. 293/huNaIII18-300-20 had an activation threshold voltage of -40 mV (Figure 9A), a steady state $V_{1/2}$ inactivation voltage of -58 mV (Figure 9B), a recovery time after inactivation of 2.5 ms (fast component) AND 113 ms (slow component-(Figure 9C), and inactivation kinetics of
30 0.8 ms (Figure 9D).

The present invention is not to be limited in scope by the specific embodiments described herein. Indeed, various modifications of the invention in addition to those described herein will become apparent to those skilled in the art from the foregoing description and the accompanying figures. Such modifications are
5 intended to fall within the scope of the appended claims.

Patents, patent applications, publications, procedures, and the like are cited throughout this application, the disclosures of which are incorporated herein by reference in their entireties.

WHAT IS CLAIMED IS:

1. An isolated nucleic acid comprising a nucleotide sequence encoding a polypeptide
having the amino acid sequence of Figure 2 (SEQ ID NO: 2).
2. The isolated nucleic acid of claim 1, comprising the nucleotide sequence of Figure 1 (SEQ ID NO: 1).
3. A recombinant vector comprising a nucleotide sequence encoding a polypeptide having the amino acid sequence of Figure 2 (SEQ ID NO:2).
4. A host cell comprising the recombinant vector of claim 3.
5. A host cell genetically engineered to comprise the nucleic acid of claim 1.
6. The host cell of claim 5 which is eukaryotic.
7. A eukaryotic host cell genetically engineered to express, or overexpress, a polypeptide having the amino acid sequence of Figure 2 (SEQ ID NO: 2).
8. A method for expressing a polypeptide in a cell cultured *in vitro* comprising culturing the cell of claim 4, 5, 6 or 7 under conditions conducive to the expression of the polypeptide comprising the amino acid sequence of Figure 2 (SEQ ID NO:2).
9. An isolated polypeptide comprising the amino acid sequence of Figure 2 (SEQ ID NO:2).

10. A host cell genetically engineered to co-express a polypeptide comprising the amino acid sequence of Figure 2 (SEQ ID NO:2) and a β -subunit of a sodium channel selected from the group consisting of $\beta 1$, $\beta 2$, and $\beta 3$.

11. An antibody or antigen-binding fragment that specifically binds to a polypeptide having the amino acid sequence of Figure 2 (SEQ ID NO: 2).

12. The antibody of claim 11, which is a monoclonal antibody.

13. A method for detecting expression in a sample of a polypeptide comprising the amino acid sequence of Figure 2 (SEQ ID NO:2), which method comprises detecting specific binding of the antibody or antigen-binding fragment of claim 11 to a polypeptide in the sample.

14. A method for identifying a test compound that binds to a sodium channel comprising a polypeptide comprising the amino acid sequence of Figure 2 (SEQ ID NO:2), which method comprises:

(i) contacting a host cell that expresses a sodium channel comprising a polypeptide comprising the amino acid sequence of Figure 2 (SEQ ID NO:2) with a test compound; and

(ii) determining whether the test compound binds to the host cell but not to a control cell that does not express a sodium channel comprising a polypeptide comprising the amino acid sequence of Figure 2 (SEQ ID NO:2).

15. An assay method for identifying a test compound that modulates the activity of a sodium channel comprising a polypeptide comprising the amino acid sequence of Figure 2 (SEQ ID NO:2), which method comprises:

(i) providing a host cell that expresses a functional sodium channel comprising at least one polypeptide comprising the amino acid sequence of Figure 2 (SEQ ID NO:2),

(ii) contacting the host cell with a test compound under conditions that would activate sodium channel activity of said functional sodium channel in the absence of

the test compound; and

(iii) determining whether the host cell contacted with the test compound exhibits a modulation in activity of the functional sodium channel.

16. The assay method of claim 15, wherein the host cell has been genetically engineered to express or overexpress the functional sodium channel.

17. The assay method of claim 15, wherein the host cell has been genetically engineered by the introduction into the cell of a nucleic acid molecule having a nucleotide sequence encoding a polypeptide comprising the amino acid sequence of Figure 2 (SEQ ID NO:2).

18. The assay method of claim 15, wherein the host cell has been genetically engineered to upregulate the expression of a nucleic acid encoding a polypeptide comprising the amino acid sequence of Figure 2 (SEQ ID NO:2),

19. The assay method of claim 18, wherein the upregulated nucleic acid is endogenous to the host cell.

20. The assay method of claim 15, wherein the modulation of the functional sodium channel activity is antagonism of that activity.

21. The assay method of claim 15, wherein the modulation of the functional sodium channel activity is agonism of that activity.

FIGURE 1: NaIIII18 cDNA (SEQ ID NO: 1)

tgaaaagatggcacaggcactgttggtagccccaggacctgaaagcttccgcctttttactaga
gaatctcttgctgctatcgaaaaacgtgctgcagaagagaaagccaagaagcccaaaaaggaac
aagataatgatgatgagaacaaaccaaagccaaatagtgacttggaagctggaaagaaccttcc
atttattttatggagacattcctccagagatgggtgtcagagcccttgaggacctggatccctac
tatatcaataagaaaactttttatagtaataaaggaaaggcaattttccgattcagtgcca
cctctgccttgctatattttaactccactaaacctgttaggaaaattgctatcaagattttgggt
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cacttataaaaaatcttggcaagaggggttttgcctagaagattttacgtttcttcgtgatccatg
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gtttcagcccttcgaactttcagagtcttgagagctctgaaaactatttctgtaattccagggtt
taaagaccattgtgggggcccctgatccagtcggtaagaagctttctgatgtgatgatcctgac
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aatgtttgcagtgcccccaagcgattctgcttttgaaaccaacaccacttccctactttaatg
gcacaatggattcaaatgggacattttgttaatgtacaatgagcacatttaactggaaggatta
cattggagatgacagtcacttttatgttttggatggggcaaaaagaccctttactctgtggaat
ggctcagatgcaggccagtgctccagaaggatacatctgtgtgaaggctggctcgaaaccccaact
atggctacacaagctttgacaccttttagctgggctttcctgtctctatttcgactcatgactca
agattactgggaaaatctttaccagttgacattacgtgctgctgggaaaacatacatgatattt
tttgtcctggctcattttcttgggctcattttatttgggtgaattttgatcctggctgtgggtggcca
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gcaccttgaaaggaaacaacaaaggagagagagacagctttcccaaattccgaatctgaagacagc
gtcaaaagaagcagcttccctttctccatggatggaaacagactgaccagtgcacaaaaattct
gctcccctcatcagctctctcttgagtatccgtggctccctgttttcccaagacgcaatagcaa
aacaagcattttcagtttcagaggtcgggcaaaggatgttggatctgaaaatgactttgctgat
gatgaacacagcacatttgaagacagcgaaagcaggagagactcactgtttgtgcccacagac
atggagagcgacgcaacagtaacgttagtcaggccagtatgtcatccaggatgggtgccagggt
tccagcaaatgggaagatgcacagcactgttgattgcaatgggtgtgggtttccttgggtgggtgga
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catttttgtgtgttggcagcagctctttggtaagagctacaaagaatgtgtctgcaagatc
aatgatgactgtacgctcccacgggtggcacatgaacgacttcttccactccttctgattgtgt

FIGURE 1 (continued)

tccgcgtgctgtgtggagagtggatagagaccatgtgggactgtatggagggtcgctggccaaac
catgtgccttattgttttcatgttgggtcatgggtcattggaaaccttggtggttctgaacctcttt
ctggccttattgttgagttcatttagctcagacaaccttgctgctactgatgatgacaatgaaa
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gagatgggaatggaaccaccagtggtgttaggtactggaagcagtggtgaaaaatacgtaatcga
tgaaatgattatatgtcattcataaacaacccagcctcaccgtcacagtgccaattgctggt
ggagagtctgactttgaaaacttaataactgaagagttcagcagtgagtcagaactagaagaaa
gcaaagagaaattaaatgcaaccagctcatctgaaggaagcacagttgatgttggttctaccccg
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atcaaaaccatgctagaatatgctgacaaagtctttacctatatattcattctggaaatgcttc
tcaaatgggttgcttatggatttcaaacatatttactaatgcctgggtgctggctagatttctt
gatcgttgatgtttctttgggttagcctggtagccaatgctcttggtactcagaactcgggtgcc
atcaaatcattacggacattaagagctttaagacctctaagagccttatcccggtttgaaggca
tgaggggtggttgatgtctcttggttgagcaattccctctatcatgaatgtgctggttggtctg
tctcatcttctggttgatcttttagcatcatgggtgtgaatttggttgctggcaagttctaccac
tgtgttaacatgacaacgggttaacatgtttgacattagtgatgttaacaatttgagtgactgtc
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gattcacgagatgttaaacttcagcctgtatatgaagaaaatctgtacatgtatttatactttg
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acaaattccaaggaatggtctttgatgtttgtaaccagacaagtctttgatatcagcatcatgat
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tcgtctccctcagacactactacttactataggctggaacatctttgactttgtggtggtgat
tctctccattgtaggtatgtttctggctgagatgatagaaaagtattttgtgtcccctaccttg
ttccgagtgatccgtcttgccaggattggccgaatcctacgtctgatcaaaggagcaaaggga
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gctggaattgatgacatgttcaactttgagacctttggcaacagcatgatctgcttggttccaaa
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agcttatttgccatggatctgccatgggtcagtggtgaccggatccactgtcttgatattttatt
tgcccttacaaagcgtgttttgggtgagagtggagagatggatgcccttcgaatacagatggaa
gacaggtttatggcatcaaacccctccaaagtctcttatgagcctattacaaccactttgaaac

FIGURE 1 (continued)

gtaaacaagaggaggtgtctgccgctatcattcagcgtaatttcagatggttatcttttaaagca
aagggttaaaaaatatatcaagtaactataacaaagaggcaattaaagggaggattgacttacct
ataaaaacaagacatgattattgacaaactaaatgggaactccactccagaaaaaacagatggga
gttcctctaccacctctcctccttcctatgatagtgtaacaaaaccagacaaggaaaagtttga
gaaagacaaaccagaaaaagaaagcaaaggaaaagaggtcagagaaaatcaaaagtaaaaagaa
acaaagaattatctttgtgatcaattgtttacagcctatga

FIGURE 2: NaIIII18 amino acid (SEQ ID NO: 2)

MAQALLVPPGPESFRLFTRESLAAIEKRAAEEKAKKPKKEQDNDNENKPKPNSDLEAG
KNLPFIYGDIPPEMVSEPLEDLDPYYINKKTFIVMNGKKAIFRFSATSALYILTPLNPVR
KIAIKILVHSLFSMLIMCTILTNCVFMTLSPDWTKNVEYTFGTGIYTFESLIKILARGF
CLEDFTFRLDPWNWLDVSVIVMAYVTEFVSLGNVSALRTFRVLRALKTISVIPGLKTIVG
ALIQSVKKLSVDMILTVFCLSVFALIGLQLFMGNLRNKCLQWPPSDSAFETNTTSYFNGT
MDSNGTFVNVTMSTFNWKDYIGDDSHFYVLDGQKDPLLCGNGSDAGQCPEGYICVKAGR
PNYGYTSFDTFSWAFSLFRLMTQDYWENLYQLTLRAAGKTYMIFVVLVIFLGSFYLVNL
ILAVVAMAYEEQNQATLEEAQKEAEFQQMLEQLKKQEEAQAVAAASAASRDFSGIGGL
GELLESSSEASKLSSKSAKEWRNRKRKRREHLEGNNKGERDSFPKSESEDSVKRSSFL
FSMDGNRLTSDKKFCSPHQSLLSIRGSLFSPRNSKTSIFSFRGRAKDVGSSENDFADEH
STFEDSESRRDSLFPVPHRHGERNSNVSQASMSSRMVPGLPANGKMHSTVDCNGVVSLVG
GPSALTSPGTGQLPPEGTTTETEVKRRLSSYQISMEMLEDSSGRQRAVSIASILTNTMEE
LEESRQKCPPCWYRFANVFLIWDCCDAWLKVKHLVNLIVMDPFVDLAITICIVLNTLFMA
MEHYPMTEQFSSVLTVGNLVFTGIFTAEMVLKIIAMDPIYFQEGWNIFDGIIVSLSLME
LGLSNVEGLSVLRSFRLRLRVFKLAKSWPTLNMLIKIIGNSVGALGNLTLVLAIIVFIFAV
VGMQLFGKSYKECVCKINDCTLPRWHMNDFFHSFLIVFRVLCGEWIETMWDCMEVAGQT
MCLIVFMLVMVIGNLVVLNLFALLSSFSNDLAATDDDNEMNNLQIAVGRMQKGIDYV
KNKMRECFQKAFFRKPKVIEIHEGNKIDSCMSNNTGIEISKELNYLRDGNNGTTSVGTGS
SVEKYVIDENDYMSFINNPSTVTVPIAVGESDFENLNTEEFSSSESELESKEKLNATSS
SEGSTVDVVLPREGEQAETEPEEDLKPEACFTEGCIKKFPFCQVSTEEGKGKIWWNLRKT
CYSIVEHNNWFETFIVFMILLSSGALAFEDIYIEQRKTIKTMLEYADKVFTYIFILEMLLK
WVAYGFQTYFTNAWCWLDFLIVDVSLVSLVANALGYSELGAIKSLRTLRLRPLRLSRF
EGMRVVVNALVGAIIPSIMNVLLVCLIFWLIFSIMGVNLFAGKFYHCVNMTTGNMFDISDV
NNLSDCQALGKQARWKNVKNFNDNVGAGYLALLOQVATFKGWMDIMYAAVDSRDVKLQPVY
EENLYMYLYFVIFIIIFGSFFTLNLFIVGVIDNFNQKKKFGGQDIFMTEEQKKYNNAMKK
LGSKKPQKPIPRPANKFQGMVDFVTRQVFDISIMILICLNMVTMMVETDDQGYMTLV
SRINLVFIVLFTGEFVLRRLVSLRHYYFTIGWNIFDFVVVILSIVGMFLAEMIEKYFVSPT
LFRVIRLARIGRILRLIKGAKGIRTLFLALMMSLPALFNIGLLFLVMFIYAI FGMSNFA
YVKKEAGIDDMFNFETFGNSMICLFQITTSAGWDGLLAPILNSAPPDCPDTHPGSSVK
GDCGNPSVGIFFFVSYIIISFLVVVNMYIAVILENFSVATEESAEPLEDDFEMFYEVWE
KFDPDATQFIEFSKLSDFAAALDPPLLIAPKNKVQLIAMDLPMVSGDRIHCLDILFAFTK
RVLGESGEMDALRIQMEDRFMASNPSKVSYPEITTTTLKRKQEEVSAAIIQRNFRCYLLKQ
RLKNISSNYNKEAIKGRIDLPKQDMIIDKLNGNSTPEKTDGSSSTTSPPSYDSVTKPKD
EKFEKDKPEKESKGKEVRENQK

**FIGURE 3: cDNA sequence of human SCN3A of Clare et al.
(SEQ ID NO: 3)**

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1  taccctaacc atcttggatg ctgggctttg ttatgctgta attcataagg ctctgtttta
61  tcagagatta tggagcaaga aaactgaagc caagccacat caaggtttga cagggatgag
121 atacctgtca aggattcata gtagagtggc ttactgggaa aggagcaaag aatctcttct
181 agggatattg taagaataaa tgagataatt cacagaaggg acctggagct tttccgaaa
241 aagggtgctgt gactatctaa ggtaattcgt atgcaagaag ctacacgtaa ttaaattgtgc
301 aggatgaaaa gatggcacag gcactgttgg taccgccagg acctgaaagc ttccgccttt
361 ttactagaga atctcttgct gctatcgaaa aacgtgctgc agaagagaaa gccaaagaagc
421 caaaaaggga acaagataat gatgatgaga acaaaccaaa gccaaatagt gacttgggaag
481 ctggaaagaa ccttccattt atttatggag acattcctcc agagatgggtg tcagagcccc
541 tggaggacct ggatccctac tatatcaata agaaaacttt tatagtaatg aataaaggaa
601 aggcaatttt ccgattcagt gccacctctg ccttgtatat tttaactcca ctaaaccctg
661 ttaggaaaaat tgctatcaag attttgggtac attctttatt cagcatgctt atcatgtgca
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781 tagagtacac attcactgga atctataacct ttgagtcact tataaaaatc ttggcaagag
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901 tcattgtgat ggcgtatgta acagaatttg taagcctagg caatgtttca gccctcgaa
961 ctttcagagt cttgagagct ctgaaaacta tttctgtaat tccaggttta aagaccattg
1021 tgggggcccc gatccagtcg gtaaagaagc tttctgatgt gatgatcctg actgtgttct
1081 gtctgagcgt gtttgctctc attgggctgc agctgttcat gggcaatctg aggaataaat
1141 gtttgagctg gcccccaagc gattctgctt ttgaaaccaa caccacttcc tactttaatg
1201 gcacaatgga ttcaaatggg acatttgtta atgtaacaat gagcacattt aactggaagg
1261 attacattgg agatgacagt cacttttatg ttttggatgg gcaaaaagac cctttactct
1321 gtggaaatgg ctcatgca ggccagtgct cagaaggata catctgtgtg aaggctgggtc
1381 gaaaccccaa ctatggctac acaagctttg acacctttag ctgggctttc ctgtctctat
1441 ttcgactcat gactcaagac tactgggaaa atctttacca gttgacatta cgtgctgtg
1501 ggaaaacata catgatattt tttgtcctgg tcattttctt gggctcattt tatttgggtga
1561 atttgatcct ggctgtggtg gccatggcct atgaggagca gaatcaggcc accttgggaag
1621 aagcagaaca aaaagaggcc gaatttcagc agatgctcga acagcttaaa aagcaacagg
1681 aagaagctca ggcagtgtcg gcagcatcag ctgcttcaag agatttcagt ggaatagggtg
1741 ggttaggaga gctgttggaa agttcttcag aagcatcaaa gttgagttcc aaaagtgtca
1801 aagaatggag gaaccgaagg aagaaaagaa gacagagaga gcaccttgaa ggaaacaaca
1861 aaggagagag agacagcttt cccaaatccg aatctgaaga cagcgtcaaa agaagcagct
1921 tccttttctc catggatgga aacagactga ccagtgaaca aaaattctgc tccctcatc
1981 agtctctctt gagtatccgt ggctccctgt tttcccaag acgcaatagc aaaacaagca
2041 ttttcagttt cagaggtcgg gcaaaggatg ttggatctga aaatgacttt gctgatgatg
2101 aacacagcac atttgaagac agcgaagca ggagagactc actgtttgtg ccgcacagac
2161 atggagagcg acgcaacagt aacggcacca ccactgaaac ggaagtcaga aagagaaggt
2221 taagctctta ccagatttca atggagatgc tggaggattc ctctggaagg caaagagccg
2281 tgagcatagc cagcattctg accaacacaa tggagaactc tgaagaatct agacagaaat
2341 gtccgccatg ctggtataga tttgccaatg tgttcttgat ctgggactgc tgtgatgcat
2401 ggttaaaagt aaaacatctt gtgaatttaa ttgttatgga tccatttgtt gatcttgcca
2461 tcactatttg cattgtctta aataccctct ttatggccat ggagcactac cccatgactg
2521 agcaattcag tagtgtgttg actgtaggaa acctggtctt tactgggatt ttcacagcag
2581 aaatggttct caagatcatt gccatggatc cttattacta tttccaagaa ggctggaata
2641 tctttgatgg aattattgtc agcctcagtt taatggagct tggctgtgca aatgtggagg
2701 gattgtctgt actgcatca ttcagactgc ttagagtttt caagttggca aaatcctggc
2761 ccacactaaa tatgctaatt aagatcattg gcaattctgt gggggctcta ggaaacctca
2821 ccttgggtgt ggccatcatc gtcttcattt ttgctgtggt cggcatgcag ctctttggta
2881 agagctacaa agaattgtgtc tgcaagatca atgatgactg tacgctccca cgggtggcaca
2941 tgaacgactt ctccactcc ttcttgattg tgttccgcgt gctgtgtgga gagtggatag
3001 agaccatgtg ggactgtatg gaggtgcgtg gccaaacctat gtgccttatt gttttcatgt
3061 tgggtcatgg cattggaaac cttgtggttc tgaacctctt tctggcctta ttgttgagtt
3121 catttagctc agacaacctt gctgctactg atgatgacaa tgaaatgaat aatctgcaga

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FIGURE 3 (continued)

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3181 ttgcagtagg aagaatgcaa aagggaattg attatgtgaa aaataagatg cgggagtggt
3241 tccaaaaagc ctttttttaga aagccaaaag ttatagaaat ccatgaaggc aataagatag
3301 acagctgcat gtccaataat actggaattg aaataagcaa agagcttaat tatcttagag
3361 atgggaatgg aaccaccagt ggtgtaggta ctggaagcag tgttgaaaaa tacgtaatcg
3421 atgaaaatga ttatatgtca ttcataaaca accccagcct caccgtcaca gtgccaatgt
3481 ctgttgaggaga gtctgacttt gaaaacttaa atactgaaga gttcagcagt gagtccagaac
3541 tagaagaaaag caaagagaaa ttaaattgcaa ccagctcatc tgaagggaagc acagttgatg
3601 ttgttctacc ccgagaaggt gaacaagctg aaactgaacc cgaagaagac cttaaaccgg
3661 aagcttggtt tactgaagga tgtattaaaa agtttccatt ctgtcaagta agtacagaag
3721 aaggcaaagg gaagatctgg tggaaatcttc gaaaaacctg ctacagtatt gttgagcaca
3781 actggtttga gactttcatt gtgttcatga tccttctcag tagtggtgca ttggcctttg
3841 aagatatata cattgaacag cgaaagacta tcaaaaccat gctagaatat gctgacaaag
3901 tctttaccta tatattcatt ctggaaatgc ttctcaaatg gggtgcttat ggatttcaaa
3961 catatttcac taatgcctgg tctggcctag atttcttgat cgttgatgtt tctttggtta
4021 gcctggtagc caatgctctt ggctactcag aactcgggtg catcaaatca ttacggacat
4081 taagagcttt aagacctcta agagccttat cccggtttga aggcattgagg gtggttgtga
4141 atgctcttgt tggagcaatt cctctatca tgaatgtgct gttggtctgt ctcatcttct
4201 gggtgatctt tagcatcatg ggtgtgaatt tgtttgctgg caagttctac cactgtgtta
4261 acatgacaac gggtaacatg tttgacatta gtgatgttaa caatttgagt gactgtcagg
4321 ctcttgccaa gcaagctcgg tggaaaaacg tgaaagtaaa ctttgataat gttggcgctg
4381 gctatcttgc actgcttcaa gtggccacat tttaaaggctg gatggatatt atgtatgcag
4441 ctgttgattc acgagatgtt aaacttcagc ctgtatatga agaaaatctg tacatgtatt
4501 tctatttgt catctttatc atctttgggt catctctcac tctgaatcta tctattggtg
4561 tcatcataga taacttcaac cagcagaaaa agaagtttgg aggtcaagac atctttatga
4621 cagaggaaca gaaaaaatat tacaatgcaa tgaagaaact tggatccaag aaacctcaga
4681 aacctatacc tcgcccagca aacaaattcc aaggaaatgg ctttgatttt gtaaccagac
4741 aagtctttga tatcagcatc atgatcctca tctgcctcaa catggtcacc atgatggtg
4801 aaacggatga ccagggcaaa tacatgacct tagttttgtc ccgatcaac ctagtgttca
4861 ttgttctgtt cactggagaa tttgtgctga agctcgtctc cctcagacac tactacttca
4921 ctataggctg gaacatcttt gactttgtgg tggatgattct ctccattgta ggtatgtttc
4981 tggctgagat gatagaaaag tattttgtgt cccctacctt gttccgagt atccgtcttg
5041 ccaggattgg ccgaatccta cgtctgacca aaggagcaaa ggggatccgc acgctgctct
5101 ttgctttgat gatgtccctt cctgcgttgt ttaacatcgg cctcctgctc ttcctgggtca
5161 tgtttatcta tgccatcttt gggatgtcca actttgccta tgttaaaaag gaagctggaa
5221 ttgatgacat gttcaacttt gagacctttg gcaacagcat gatctgcttg ttccaaatta
5281 caacctctgc tggctgggat ggattgctag cacctattct taatagtgc ccacccgact
5341 gtgacctga cacaattcac cctggcagct cagttaaggg agactgtggg aacctatctg
5401 ttgggatttt ctttttcgtc agttacatca tcatatcctt cctggttgtg gtgaacatgt
5461 acatcgcggt catcctggag aacttcagtg ttgctactga agaaagtgc gagcccctga
5521 gtgaggatga ctttgagatg ttctatgagg tttgggaaaa gtttgatccc gatgagcccc
5581 agtttataga gttctctaaa ctctctgatt ttgcagctgc cctggatcct cctcttctca
5641 tagcaaaacc caacaaagtc cagcttattg ccatggatct gccatgggtc agtggtgacc
5701 ggatccactg tcttgatatt ttatttgcct ttacaaagcg tgttttgggt gagagtggag
5761 agatggatgc ccttcgaata cagatggaag acaggtttat ggcacaaac cctccaaag
5821 tctcttatga gcctattaca accactttga aacgtaaaaca agaggaggtg tctgcogcta
5881 tcattcagcg taatttcaga tgttatcttt taaagcaaag gttaaaaaat atatcaagta
5941 actataacaa agaggcaatt aaagggagga ttgacttacc tataaaacaa gacatgatta
6001 ttgacaaact aaatgggaac tccactccag aaaaaacaga tgggagttcc tctaccacct
6061 ctctctcttc ctatgatagt gtaacaaaac cagacaagga aaagtttgag aaagacaaac
6121 ccagaaaaga aagcaaagga aaagaggtca gagaaaatca aaagtaaaaa gaaacaaaga
6181 attatctttg tgatcaattg tttacagcct atgaaggtaa agtatatgtg tcaactggac
6241 ttcaagagga ggtccatgcc aaactgactg ttttaacaaa tactcatagt cagtgcctat
6301 acaagacagt gaagtgcct ctctgtcact gcaactctgt gaagcagggt atcaacattg
6361 acaagaggtt gctgttttta ttaccagctg acactgctga ggagaaaccc aatggctacc

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FIGURE 3 (continued)

6421 tagactatag ggatagttgt gcaaagtgaa cattgtaact acaccaaaca ccttttagtac
6481 agtccttgca tccattctat ttttaacttc catatctgcc atatTTTTac aaaatttggt
6541 ctagtgcatt tccatgggcc ccaattcata gtttattcat aatgctatgt cactatTTT

FIGURE 4: amino acid sequence of human SCN3A (SEQ ID NO: 4)

MAQALLVPPGPESFRLFTRESLAAIEKRAAEKAKKPKKEQDNDDENKPKPNSDLEAGKNLPFI
YGDIPPEMVSEPLEDLDPYYINKKTFIVMNKGKAI FRFSATSALYILTPLNPVRKIAIKILVHS
LFSMLIMCTILTNCVFMTLSNPPDWTKNVEYTFGTGIYTFESLIKILARGFCLEDFTLRDPWNW
LDFSIVIMAYVTEFVSLGNVSALRTFRVLRALKTISVIPGLKTIVGALIQSVKKLSVDMILTVF
CLSVFALIGLQLFMGNLRNKCLQWPPSDSAFETNTTSYFNGTMDSNGTFVNVMTSTFNWKDYIG
DDSHFYVLDGQKDPLL CGNGSDAGQCPEGYICVKAGRNPNYGYTSFDTFSWAFLSLFRMTQDY
WENLYQLTLRAAGKTYMIFVVLVIFLGSFYLVNLILAVVAMAYEEQNQATLEEAEQKEAEFQQM
LEQLKKQQEEAQAVAAASAASRDFSGIGGLGELLESSSEASKLSSKSAKEWRNRKRQRREHL
EGNNKGERDSFPKSESEDSVKRSSFLFSMDGNRLTSDKKFCSPHQSLLSIRGSLFSPRRNSKTS
IFSFRGRAKDVGSENFADDEHSTFEDSESRRDSLFPVPHRHGERRNSNGTTTETEVKRRLSSY
QISMEMLEDSSGRQRAVSIA SILTNTMEELEESRQKCPPCWYRFANVFLIWDCCDAWLKVKHLV
NLIVMDPFVDLAIITICIVLNTLFMAMEHYPMTEQFSSVLTVGNLVFTGIFTAEMVLKIIAMDPY
YYFQEGWNI F DGIIVSLSLMELGLSNVEGLSVLRSFRLLRVFKLAKSWPTLNMLIKIIGNSVGA
LGNLTLLVLAII VFIFAVVGMQLFGKSYKECVCKINDDCTLPRWHMNDFFHSFLIVFRVLCGEWI
ETMWDCMEVAGQTMCLIVFMLVMVIGNLVVLNLFALLSSSFSSDNLAATDDD NEMNNLQI AVG
RMQKGIDYVKNKMRECFQKAFFRKPKVIEIHEGNKIDSCMSNNTGIEISKELNYLRDNGTTSG
VGTGSSVEKYVIDENDYMSFINNPSLTVTVPIAVGESDFENLNTEEFSSSESELESKEKLNATS
SSEGSTVDVVLPREGEQAETEPEEDLKPEACFTEGCIKKFPFCQVSTEEGKGKIWWNLRKTCYS
IVEHWNWFETFIVFMILLSSGALAFEDIYIEQRKTIKTMLEYADKVFTYIFILEMLLKWVAYGFQ
TYFTNAWCWLD FLIVDVSLVSLVANALGYSELGAIKSLRTLRLALRPLRALS RFEGMRVVNALV
GAIPSIMNVLLVCLIFWLI FSIMGVNLFAGKFYHCVNMTTGNMFDISDVNNLSDCQALGKQARW
KNVKVNFDNVGAGYLALLQVATFKGWMDIMYAAVDSRDVKLQPVYEENLYMYLYFVIFIIFGSF
FTLNLFIGVIIDNFNQKKKFGGQDIFMTEEQKKYYNAMKKLGSKKPQKPIPRPANKFQGMVFD
FVTRQVFDISIMILICLNMVTMMVETDDQGYMTLVLSRINLVFIVLFTGEFVLKLVSLRHYYF
TIGWNI FDFVVVILSIVGMFLAEMIEKYFVSPTLFRVIRLARIGRILRLIKGAKGIRTLLFALM
MSLPALFNIGLLFLVFMFIYAI FGMSNFAYVKKEAGIDDMFNFFETFGNSMICLFQITTSAGWDG
LLAPILNSAPPD CDPD TIHPGSSVKGDCGNPSVGIFFFVSYIIISFLVVVNMYIAVILENFSVA
TEESAEPLESDDFEMFYEVWEKFDPDATQFIEFSKLSDFAAALDPPLLI AKPNKVQLIAMDLPM
VSGDRIHCLDILFAFTKRVLGESGEMDALRIQMEDRFMASNPSKVS YEPI TTTTLKRKQEEVSAA
IIQRNFRCYLLKQRLKNISSNYNKEAIKGRIDLP IKQDMIIDKLNGNSTPEKTDGSSSTTSPPS
YDSVTKPDKEKFEKDKPEKESKGKEVRENQK

FIGURE 5: cDNA of human sodium channel α -subunit variant by Jeong et al. (SEQ ID NO: 5)

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1 agcgaagcgg aggcataagc agagaggatt ctggaaaggt ctctttgttt tcttatccac
61 agagaaagaa agaaaaaaaaa ttgtaactaa tttgtaaacc tctgtggtca aaaaaaaaaa
121 aaaaaaaaaa gctgaacagc tgccagagga agacacgtta taccctaacc atcttggatg
181 ctgggctttg ttatgctgta attcataagg ctctgtttta tcagagatta tggagcaaga
241 aaactgaagc caagccacat caaggtttga cagggatgag atacctgtca aggattcata
301 gtagagtggc ttactgggaa aggagcaaag aatctcttct agggatattg taagaataaa
361 tgagataaatt cacagaaggg acctggagct tttccggaaa aagggtgctgt gactatctaa
421 ggtaattcgt atgcaagaag ctacacgtaa ttaaatgtgc aggatgaaaa gatggcacag
481 gcactgttgg tacccccagg acctgaaagc ttccgccttt ttactagaga atctcttgct
541 gctatcgaaa aacgtgctgc agaagagaaa gccaaagaagc ccaaaaagga acaagataat
601 gatgatgaga acaaaccaaa gccaaatagt gacttggaag ctggaaagaa ccttccattt
661 atttatggag acattcctcc agagatggtg tcagagcccc tggaggacct ggatccctac
721 tatatcaata agaaaacttt tatagtaatg aataaaggaa aggcaatttt ccgattcagt
781 gccacctctg ccttgtatat ttttaactcca ctaaaccctg ttaggaaaat tgctatcaag
841 attttgggtac attcttttatt cagcatgctt atcatgtgca ctattttgac caactgtgta
901 ttttatgacct tgagcaacct tcctgactgg acaaagaatg tagagtacac attcactgga
961 atctataacct ttgagtcact tataaaaatc ttggcaagag ggttttgctt agaagatttt
1021 acgtttcttc gtgatccatg gaactggctg gatttcagtg tcattgtgat ggcatatgtg
1081 acagagtttg tggacctggg caatgtctca gcgttgagaa cattcagagt tctccgagca
1141 ctgaaaacaa tttcagtcac tccagggttta aagaccattg tgggggccct gatccagtcg
1201 gtaaagaagc tttctgatgt gatgatcctg actgtgttct gtctgagcgt gtttgctctc
1261 attgggctgc agctgttcat gggcaatctg aggaataaat gtttgagtg gcccccaagc
1321 gattctgctt ttgaaaccaa caccacttcc tactttaatg gcacaatgga ttcaaatggg
1381 acatttgttt atgtaacaat gagcacattt aactggaagg attacattgg agatgacagt
1441 cacttttatg ttttggatgg gcaaaaagac ctttactct gtggaaattg ctcatgca
1501 ggccagtgtc cagaaggata catctgtgtg aaggctgggtc gaaaccccaa ctatggctac
1561 acaagctttg acacctttag ctgggctttc ctgtctctat ttcgactcat gactcaagac
1621 tattgggaaa atctttacca gttgacatta cgtgctgctg ggaaaacata catgatattt
1681 tttgtcctgg tcatttttctt gggctcattt tatttgggtg atttgatcct ggctgtgggtg
1741 gccatggcct atgaggagca gaatcaggcc accttggaag aagcagaaca aaaagaggcc
1801 gaatttcagc agatgctcga acagcttaaa aagcaacagg aagaagctca ggcagttgctg
1861 gcagcatcag ctgcttcaag agatttcagt ggagtaggtg ggttaggaga gctgttggaa
1921 agttcttcag aagcatcaaa gttgagttcc aaagggtgcta aagaatggag gaaccggagg
1981 aagaaaagaa gacagagaga gcaccttgaa ggaaacaaca aaggagagag agacagcttt
2041 cccaaatccg aatctgaaga cagcgtcaaa agaagcagct tcctttctc cctggatgga
2101 aacagactga ccagtgacaa aaaatctctg tccccctcat agtctctctt gagtatccgt
2161 ggctccctgt tttccccaag acgcaatagc aaaacaagca ttttcagttt cagaggtcgg
2221 gcaaaggatg ttggatctga aaatgacttt gctgatgatg aacacagcac atttgaagac
2281 ggcgaaagca ggagagactc actgtttgtg ccgcacagac atggagagcg acgcaacagt
2341 aacgttagtc aggccagtat gtcacccagg atgggtgccag ggcttccagc aaatgggaag
2401 atgcacagca ctgtggattg caatgggtgtg gtttccttgg tgggtggacc ttcagctcta
2461 acgtcaccta ctggacaact tccccagag ggcaccacca ctgaaacgga agtcagaaag
2521 agaaggttaa gctcttacca gatttcaatg gagatgctgg aggattcctc tggaaaggcaa
2581 agagccgtga gcatagccag cattctgacc aacacaatgg aagaacttga agaacttaga
2641 cagaaatgtc cgccatgctg gtatagattt gccaatgtgt tcttgatctg ggactgctgt
2701 gatgcatggg taaaagtaaa acatcttgtg aatttaattg ttatggatcc atttgttgat
2761 cttgccatca ctatttgcat tgtcttaaat accctcttta tggccatgga gcactacccc
2821 atgactgagc aattcagtag tgtgttgact gtaggaaacc tggctcttac tgggattttc
2881 acagcagaaa tggttctcaa gatcattgcc atggatcctt attactattt ccaagaaggc
2941 tggaaatatc ttgatggaat tattgtcagc ctcaagttta tggagcttgg tctgtcaaat
3001 gtggagggat tgtctgtact gcgatcattc agactgctta gagttttcaa gttggcaaaa
3061 tcctggccca cactaaatat gctaattaa atcattggca attctgtggg ggctctagga
3121 aacctcacct tgggtgttggc catcatcgct ttcatttttg ctgtggctcg catgcagctc

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FIGURE 5 (continued)

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3181 tttggtaaga gctacaaaga atgtgtctgc aagatcaatg atgactgtac gctcccacgg
3241 tggcacatga acgacttctt ccactccttc ctgatttgtt tccgcgtgct gtgtggagag
3301 tggatagaga ccatgtggga ctgtatggag gtcgctggcc aaaccatgtg ccttattgtt
3361 ttcattgttg tcatggtcac tggaaacctt gtggttctga acctcttctt ggccttatta
3421 ttgagttcat ttagctcaga caaccttgct gctactgatg atgacaatga aatgaataat
3481 ctgcagattg cagtaggaag aatgcaaaag ggaattgatt atgtgaaaaa taagatgcgg
3541 gagtgtttcc aaaaagcctt ttttagaaag ccaaaagtta tagaaatcca tgaaggcaat
3601 aagatagaca gctgcatgtc caataatact ggaattgaaa taagcaaaga gcttaattat
3661 cttagagatg ggaatggaac caccagtggg taggtactg gaagcagtgt tgaaaaatac
3721 gtaatcgatg aaaatgatta tatgtcattc ataaacaacc ccagcctcac cgtcacagtg
3781 ccaattgctg ttggagagtc tgactttgaa aacttaaata ctgaagagtt cagcagtga
3841 tcagaactag aagaaagcaa agagaaatta aatgcaacca gctcatctga aggaagcaca
3901 gttgatgttg ttctaccccg agaaggtgaa caagctgaaa ctgaaccgga agaagacttt
3961 aaaccggaag cttgttttac tgaagggtgt attaaaaagt ttccattctg tcaagtaagt
4021 acagaagaag gcaaagggaa gatctgggtg aatcttcgaa aaacctgcta cagtattgtt
4081 gagcacaact ggtttgagac ttctattgtg ttcatgatcc ttctcagtag tgggtgcattg
4141 gcctttgaag atatatacat tgaacagcga aagactatca aaaccatgct agaatatgct
4201 gacaaagtct ttacctatat attcattctg gaaatgcttc tcaaatgggt tgcttatgga
4261 tttcaaacat atttcaactaa tgcttgggtg tggttagatt tcttgatcgt tgatgtttct
4321 ttggttagcc tggtagccaa tgctcttggc tactcagaac tcggtgccat caaatcatta
4381 cggacattaa gagctttaag acctctaaga gccttatccc ggtttgaagg catgaggggtg
4441 gttgtgaatg ctcttgttgg agcaattccc tctatcatga atgtgtgtct ggtctgtctc
4501 atcttctggg tgatcttttag catcatgggt gtgaatttgt ttgctggcaa gttctaccac
4561 tgtgttaaca tgacaacggg taacatgttt gacattagtg atgttaacaa tttgagtga
4621 tgtcaggctc ttggcaagca agctcgggtg aaaaacgtga aagtaaactt tgataatgtt
4681 ggcgctggct atcttgcact gcttcaagtg gccacattta aaggctggat ggatattatg
4741 tatgcagctg ttgattcacg agatgttaaa cttcagcctg tatatgaaga aaatctgtac
4801 atgtatttat actttgtcat ctttatcatc tttgggtcat tcttcaactc gaatctattc
4861 attggtgtca tcatagataa cttcaaccag cagaaaaaga agtttggagg tcaagacatc
4921 tttatgacag aggaacagaa aaaatattac aatgcaatga agaaacttgg atccaagaaa
4981 cctcagaaac ccatacctcg ccagcaaac aaattccaag gaatggtctt tgattttgta
5041 accagacaag tctttgatag cagcatcatg atcctcatct gcctcaacat ggtcaccatg
5101 atggtggaaa cggatgacca gggcaaatat atgaccctag ttttgtcccg gatcaacctt
5161 gtgttcattg ttctgttcac tggagaattt gtgctgaagc tcgtttccct cagacactac
5221 tacttcaacta taggctggaa catctttgac tttgtggtgg tgattctctc cattgtagggt
5281 atgtttctgg ctgagatgat agaaaagtat tctgtgtccc ctaccttgtt ccgagtgatc
5341 cgtcttgcca ggattggccg aatcctacgt ctgatcaaag gagcaaaggg gatccgcacg
5401 ctgctctttg ctttgatgat gtcccttctt gcgttgttta acatcggcct cctgctcttc
5461 ctggtcatgt ttatctatgc catctttggg atgtccaact ttgcctatgt taaaaaggaa
5521 gctggaattg atgacatggt caactttgag acctttggca acagcatgat ctgcttgttc
5581 caaattacaa cctctgctgg ctgggatgga ttgctagcac ctattcttaa tagtgcacca
5641 cccgactgtg accctgacac aattcacctt ggcagctcag ttaagggaga ccgtggggac
5701 ccatctgttg ggattttctt tttgtcagt tacatcatca tatccttctt ggttgtggtg
5761 aacatgtaca tcgcggtcat cctggagaac ttcagtgttg ctactgaaga aagtgcagag
5821 cccctgagtg aggatgactt tgagatgttc tatgaggttt gggaaaagtt tgatcccgat
5881 gcgaccaggt ttatagagtt ctctaaaact tctgattttg cagctgccct ggatcctcct
5941 cttctcatag caaaaaccaa caaagtccag cttattgcca tggatctgcc catggtcagt
6001 ggtgaccgga tccactgtct tgatatttta tttgccttta caaagcgtgt tttgtgtgag
6061 agtgagaga tggatgccct tcgaatacag atggaagaca ggtttatggc atcaaaaccc
6121 tccaaagtct cttatgagcc tattacaacc actttgaaac gtaaacaaaga ggaggtgtct
6181 gccgctatca ttcagcgtaa tttcagatgt tatcttttaa agcaaaggtt aaaaaatata
6241 tcaagtaact ataacaaaga ggcaattaaa gggaggattg acttacctat aaaacaagac
6301 atgattattg acaaactaaa tgggaactcc actccagaaa aaacagatgg gagttcctct
6361 accaccctc ctcttctcta tgatagtgtg acaaaaccag acaaggaaaa gtttgagaaa

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FIGURE 5 (continued)

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6421 gacaaaccag aaaaagaaag caaaggaaaa gaggtcagag aaaatcaaaa gtaaaaagaa
6481 acaaaagaatt atctttgtga tcaattgttt acagcctatg aaggtaaagt atatgtgtca
6541 actggacttc aagaggaggt ccatgccaaa ctgactgttt taacaaatac tcatagtcag
6601 tgcctatata agacagtga gtagacctctc tgtcactgca actctgtgaa gcagggtatc
6661 aacgttgaca agaggttgct gtttttatta ccagctgaca ctgctgagga gaaacccaat
6721 ggctacctag actatagga tagttgtgca aagtgaacat tgtaactaca ccaaaccctt
6781 ttagtacagt ccttgcattcc attctatttt taacttccat atctgccata tttttacaaa
6841 atttgttcta gtgcatttcc atgggtccca attcatagtt tattcataat gctatgtcac
6901 tattttttgt aatgaggttt acgttgaaga aacagtatac aagaaccctg tctctcaaat
6961 gatcagacaa aggtgttttt ccagagagat aaaatttttt ctcaaaacca gaaaaagaat
7021 tgtaatggct acagtttcag ttacttccat tttctagatg gctttaattt tgaaagtatt
7081 ttagtctgtt atgtttgttt ctatctgaac agttatgtgc ctgtaaaagtc tctcttaata
7141 tttaaaggat tatttttatg caaagtattc tgtttcagca agtgcaaatt ttattctaag
7201 tttcagagct ctatatttta tttagggtcaa atgctttcca aaaagtaatc taataaatcc
7261 attctagaaa aatatatcta aagtattgct ttagaatagt tgttccactt tctgctgcag
7321 tattgctttg ccatctctct ctctcagcaa agctgatagt ctatgtcaat taaataccct
7381 atgttatgta aatagttatt ttatcctgtg gtgcatgttt gggcaaatat atatatagcc
7441 tgataaaciaa cttctattaa atcaaatatg taccacagtg tatgtgtctt ttgcaagctt
7501 ccaacaggga tgtatcctgt atcattcatt aaacatagtt taaaggctat cactaatgca
7561 tgtaaatatt gcctatgctg ctctatttta ctcaatccat tcttcacaag tcttggttaa
7621 agaatgtcac atattggtga tagaatgaat tcaacctgct ctgtccatta tgtcaagcag
7681 aataatttga agctatttac aaacaccttt acttttgca ttttaattca acatgagtat
7741 catatgggat ctctctggat ttcaaggaaa cacactggat actgcctact gacaaaacct
7801 attcttcata ttttgctaaa aatatgtcta aaacttgttt aaatataaat aatgtaaaaa
7861 tataatcaac tttatttgtc agcattttgt acataagaaa attattttca ggttgatgac
7921 atcacaattt attttacttt atgcttttgc ttttgatttt taatcacaat tccaaacttt
7981 tgaatccata agatttttca atggataatt tcctaaaaata aaagttagat aatgggtttt
8041 atggatttct ttgttataat atattttcta ccattccaat aggagatata ttggtcaaac
8101 actcaaacct agatcatttt ctaccaacta tgggtgcctc aatataacct tttattcata
8161 gatgtttttt tttattcaac ttttgtagta tttacgtatg cagactagtc ttattttttt
8221 aattcctgct gcactaaagc tattacaaat ataacatgga ctttgttctt tttagccatg
8281 aacaaagtgg caaagtgttg caattaccta acatgatata aatttttgtt ttttgcaaa
8341 accaaaagtt taatgttaat tctttttaca aaactattta ctgtagtgtg ttgaagaact
8401 gcatgcaggg aattgctatt gctaaaaaga atgggtgagct acgtcattat tgagccaaaa
8461 gaataaattt cattttttat tgcatttcac ttattgggct ctgggggttt ttgtttttgt
8521 tttttgctgt tggcagttta aaatatatat aattaataaa acctgtgctt gatctgacat
8581 ttgtatacat aaaagtttac atgaatttta caacaaaacta gtgcatgatt caccaagcag
8641 tactacagaa caaaggcaaa ttaaaagcag ctttgtgaac ttttatgtgt gcaaaggatc
8701 aagttcacat gttccaactt tcagggttga taataatagt agtaaccacc tacaatagct
8761 ttcaatttca attaacctcc ttggctataa gcattctaaac tcatcttctt tcaatataat
8821 tgatgctatc tcctaattac ttgggtggcta ataaatgtta cattctttgt tacttaaatg
8881 cattatataa actcctatgt atacataagg tattaatgat atagttattg agaatttata
8941 ttaacttttt tttcaagaac ccttggtatt atgtgaggtc aaaaccaaac tcttattctc
9001 agtggaaaac tccagttgta atgcataatt ttaaagacaa tttggatcta aatatgtatt
9061 tcataattct cccataataa attatataag gtggaaaaaa aaaaaaaaaa aaaaaaaaaa
9121 aaa

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FIGURE 6: amino acid sequence of human sodium channel α -subunit variant by Jeong et al. (SEQ ID NO: 6)

MAQALLVPPGPESFRLFTRESLAAIEKRAAEKAKKPKKEQDNDDENKPKPNSDLEAGKNLPFI
 YGDIPPEMVSEPLEDLDPYYINKKTFIVMNKGKAI FRFSATSALYILTPLNPVRKIAIKILVHS
 LFSMLIMCTILTNCVFM TLSNPPDWTKNVEYFTFTGIYTFESLIKILARGFCLEDFTLRDPWNW
 LDFSIVIMAYVTEFVDLGNVSALRTRFVLRALKTISVIPGLKTI VGALIQSVKKLS DVMILTVF
 CLSVFALIGLQLFMGNLRNKCLQWPPSDSAFETNTTSYFNGTMD SNGTFFVNVTMSTFNWKDYIG
 DDSHFYVLDGQKDP LLCNGSGDAGQCPEGYICVKAGRNPNGYTSFDTFSWAFSLSLRLMTQDY
 WENLYQLTLRAAGKTYMIFVFLVIFLGSFYLVNLILAVVAMAYEEQNQATLEEAEQKEAEFQQM
 LEQLKKQQEEAQAVAAASAASRDFSGVGGGLGELLESSSEASKLSSKGAKEWNRNRKRRQREHL
 EGNNKGERDSFPKSESEDSVKRSSFLFSMDGNRLTSDKKFCSPHQSLLSIRGSLFSPRRNSKTS
 IFSFRGRAKDVGSENFADDEHSTFEDGESRRDSL FVPHRHGERRNSNVSQASMSSRMVPG LPA
 NGKMHS TVDCNGVVS LVGGPSALTSPTGQLPPEGTTTETEVKRRLSSYQISMEMLEDSSGRQR
 AVSIASILTNTMEELEESRQKCPPCWYRFANVFLIWDCCDAWLKV KHLVNLIVMDPFVDLAITI
 CIVLNTLFMAMEHYPMTEQFSSVLT VGNLVFTGIFTAEMVLKI IAMDPYFFFQEGWNIFDGIIV
 SLSLMELGLSNVEGLSVLRSFRLRLRVFKLAKSWPTLNMLIKI IGNSVGALGNLT LVLAIIVFIF
 AVVGMQLFGKSYKECVCKINDDCTLPRWHMNDFFHSFLIVFRVLCGEWIETMWDCMEVAGQ TMC
 LIVFMLVMVIGNLVVLNLF LALLLSFSSDNLAATDDDNEMNNLQIAVGRMQKGIDYVKNKMRE
 CFQKAFFRKPKVIEIHEGNKIDSCMSNNTGIEISKELNYLRDGN GTTSGVGTGSSVEKYVIDEN
 DYMSFINNPSLT VTPVPIAVGESDFENLNT EEFSSSESELEESKEKLNATSSSEGSTVDVVL PREG
 EQAETEPEEDFKPEACFTEGCIKKFPFCQVSTEEGKGKIWNLRKTCYSIVEHNWFET FIVFMI
 LLSSGALAFEDIYIEQRKTIKTMLEYADKVFTYI FILEMLLKWVAYGFQTYFTNAWCWLD FLIV
 DVSLVSLVANALGYSELGAIKSLRTLRLALRPLRALS RFEGMRVVVNALVGAIP SIMNVLLVCLI
 FWLI F SIMGVNLFAGKFYHCVNMTTG NMFDISDVNNLSDCQALGKQARWK NVKNFNDNVGAGYL
 ALLQVATFKGWMDIMYAAVDSRDVKLQPVYEENLYMYLYFVIF IIFGSFFT LNLFIGV I IDNFN
 QQKKKFGGQDI FMTEEQKKYYNAMKKLGSKKPQKPI PRPANKFQGMVFD FVTRQVFDISIMILI
 CLNMVTMMVETDDQGYMTLVLSRINLVFIVLFTGEFVLKLVSLRHYYFTIGWNIFDFV VVILS
 IVGMFLAEMIEKYSVSPTLFRVIRLARIGRILRLIKGAKGIR TLLFALMMSLPALFNIGLLLFL
 VMFIYAI FGMSNFAYVKKEAGIDDMFNFETFGNSMICLFQITTSAGWDGLLAPIILNSAPPDCDP
 DTIHPGSSSVKGDRGDPSVGIFFFVS YIIISFLVVVNMYIAVILENFSVATEESA EPLSEDDFEM
 FYEVWEKFPDPATQFIEFSKLSDFAAALDPPLLI AKPNKVQLIAMDLPMVSGDRIHCLDILFAF
 TKRVLCESGEMDALRIQMEDRFMASNPSKVS YEPITTT LKRKQEEVSAAI IQRNFRCYLLKQRL
 KNISSNYNKEAIKGRIDLP IKQDMIIDKLNGNSTPEKTDGSSSTTPPPSYDSVTKPDKEKFEKD
 KPEKESKGKEVRENQK

		Section 1				
		(1) 1	10	20	30	48
ClareAJ251507	(1)	-----	-----	-----	-----	-----
huNall18 (AK)	(1)	-----	-----	-----	-----	-----
JeongAF225987	(1)	AGCGAAGCGGAGGCATAAGCAGAGAGGATTCTGGAAAGGTCTCTTTGT				
Consensus	(1)					
		Section 2				
		(49) 49	60	70	80	96
ClareAJ251507	(1)	-----	-----	-----	-----	-----
huNall18 (AK)	(1)	-----	-----	-----	-----	-----
JeongAF225987	(49)	TTTCTTATCCACAGAGAAAGAAAGAAAAAAATTGTAACATAATTTGTA				
Consensus	(49)					
		Section 3				
		(97) 97	110	120	130	144
ClareAJ251507	(1)	-----	-----	-----	-----	-----
huNall18 (AK)	(1)	-----	-----	-----	-----	-----
JeongAF225987	(97)	AACCTCTGTGGTCAAAAAAAAAAAAAAAAAAAAAAGCTGAACAGCTGCC				
Consensus	(97)					
		Section 4				
		(145) 145	150	160	170	180
ClareAJ251507	(1)	-----	-----	-----	-----	-----
huNall18 (AK)	(1)	-----	-----	-----	-----	-----
JeongAF225987	(145)	AGAGGAAGACACGTTATACCCCTAACCATCTTGGATGCTGGGCTTTGTT				
Consensus	(145)	TACCCTAACCATCTTGGATGCTGGGCTTTGTT				
		Section 5				
		(193) 193	200	210	220	230
ClareAJ251507	(33)	ATGCTGTAATTTCATAAAGGCTCTGTTTTATCAGAGATTATGGAGCAAGA				
huNall18 (AK)	(1)	-----	-----	-----	-----	-----
JeongAF225987	(193)	ATGCTGTAATTTCATAAAGGCTCTGTTTTATCAGAGATTATGGAGCAAGA				
Consensus	(193)	ATGCTGTAATTTCATAAAGGCTCTGTTTTATCAGAGATTATGGAGCAAGA				
		Section 6				
		(241) 241	250	260	270	288
ClareAJ251507	(81)	AAACTGAAGCCAAGCCACATCAAGGTTTGGACAGGGATGAGATACCTGT				
huNall18 (AK)	(1)	-----	-----	-----	-----	-----
JeongAF225987	(241)	AAACTGAAGCCAAGCCACATCAAGGTTTGGACAGGGATGAGATACCTGT				
Consensus	(241)	AAACTGAAGCCAAGCCACATCAAGGTTTGGACAGGGATGAGATACCTGT				
		Section 7				
		(289) 289	300	310	320	336
ClareAJ251507	(129)	CAAGGATTTCATAGTAGAGTGGCTTACTGGGAAAGGAGCAAAGAATCTC				
huNall18 (AK)	(1)	-----	-----	-----	-----	-----
JeongAF225987	(289)	CAAGGATTTCATAGTAGAGTGGCTTACTGGGAAAGGAGCAAAGAATCTC				
Consensus	(289)	CAAGGATTTCATAGTAGAGTGGCTTACTGGGAAAGGAGCAAAGAATCTC				

Section 8					
	(337)	337	350	360	370 384
ClareAJ251507	(177)	TTCTAGGGGATATTGTAAGAATAAATGAGATAATTCACAGAAGGGACCT			
huNall18 (AK)	(1)	-----			
JeongAF225987	(337)	TTCTAGGGGATATTGTAAGAATAAATGAGATAATTCACAGAAGGGACCT			
Consensus	(337)	TTCTAGGGGATATTGTAAGAATAAATGAGATAATTCACAGAAGGGACCT			
Section 9					
	(385)	385	390	400	410 420 432
ClareAJ251507	(225)	GGAGCTTTTCCGGAAAAAGGTGCTGTGACTATCTAAGGTAATTTCGTAT			
huNall18 (AK)	(1)	-----			
JeongAF225987	(385)	GGAGCTTTTCCGGAAAAAGGTGCTGTGACTATCTAAGGTAATTTCGTAT			
Consensus	(385)	GGAGCTTTTCCGGAAAAAGGTGCTGTGACTATCTAAGGTAATTTCGTAT			
Section 10					
	(433)	433	440	450	460 470 480
ClareAJ251507	(273)	GCAAGAAGCTACACGTAATTAAATGTGCAGGA TGAAAAGATGGCACAG			
huNall18 (AK)	(1)	-----TGAAAAGATGGCACAG			
JeongAF225987	(433)	GCAAGAAGCTACACGTAATTAAATGTGCAGGA TGAAAAGATGGCACAG			
Consensus	(433)	GCAAGAAGCTACACGTAATTAAATGTGCAGGATGAAAAGATGGCACAG			
Section 11					
	(481)	481	490	500	510 528
ClareAJ251507	(321)	GCACTGTTGGTACCCCCAGGACCTGAAAGCTTCCGCCTTTTTTACTAGA			
huNall18 (AK)	(17)	GCACTGTTGGTACCCCCAGGACCTGAAAGCTTCCGCCTTTTTTACTAGA			
JeongAF225987	(481)	GCACTGTTGGTACCCCCAGGACCTGAAAGCTTCCGCCTTTTTTACTAGA			
Consensus	(481)	GCACTGTTGGTACCCCCAGGACCTGAAAGCTTCCGCCTTTTTTACTAGA			
Section 12					
	(529)	529	540	550	560 576
ClareAJ251507	(369)	GAATCTCTTGCTGCTATCGAAAAACGTGCTGCAGAAGAGAAAGCCAAG			
huNall18 (AK)	(65)	GAATCTCTTGCTGCTATCGAAAAACGTGCTGCAGAAGAGAAAGCCAAG			
JeongAF225987	(529)	GAATCTCTTGCTGCTATCGAAAAACGTGCTGCAGAAGAGAAAGCCAAG			
Consensus	(529)	GAATCTCTTGCTGCTATCGAAAAACGTGCTGCAGAAGAGAAAGCCAAG			
Section 13					
	(577)	577	590	600	610 624
ClareAJ251507	(417)	AAGCCCCAAAAAGGAACAAGATAATGATGATGAGAACAACCAAAGCCA			
huNall18 (AK)	(113)	AAGCCCCAAAAAGGAACAAGATAATGATGATGAGAACAACCAAAGCCA			
JeongAF225987	(577)	AAGCCCCAAAAAGGAACAAGATAATGATGATGAGAACAACCAAAGCCA			
Consensus	(577)	AAGCCCCAAAAAGGAACAAGATAATGATGATGAGAACAACCAAAGCCA			
Section 14					
	(625)	625	630	640	650 660 672
ClareAJ251507	(465)	AATAGTGACTTGGAAGCTGGAAAGAACCTTCCATTTATTTATGGAGAC			
huNall18 (AK)	(161)	AATAGTGACTTGGAAGCTGGAAAGAACCTTCCATTTATTTATGGAGAC			
JeongAF225987	(625)	AATAGTGACTTGGAAGCTGGAAAGAACCTTCCATTTATTTATGGAGAC			
Consensus	(625)	AATAGTGACTTGGAAGCTGGAAAGAACCTTCCATTTATTTATGGAGAC			

							Section 15
	(673)	673	680	690	700	710	720
ClareAJ251507	(513)	ATTCCCTCCAGAGATGGTGTCTCAGAGCCCCCTGGAGGACCTGGATCCCTAC					
huNall18 (AK)	(209)	ATTCCCTCCAGAGATGGTGTCTCAGAGCCCCCTGGAGGACCTGGATCCCTAC					
JeongAF225987	(673)	ATTCCCTCCAGAGATGGTGTCTCAGAGCCCCCTGGAGGACCTGGATCCCTAC					
Consensus	(673)	ATTCCCTCCAGAGATGGTGTCTCAGAGCCCCCTGGAGGACCTGGATCCCTAC					
							Section 16
	(721)	721	730	740	750		768
ClareAJ251507	(561)	TATATCAATAAGAAAACCTTTTATAGTAATGAATAAAGGAAAGGCAATT					
huNall18 (AK)	(257)	TATATCAATAAGAAAACCTTTTATAGTAATGAATAAAGGAAAGGCAATT					
JeongAF225987	(721)	TATATCAATAAGAAAACCTTTTATAGTAATGAATAAAGGAAAGGCAATT					
Consensus	(721)	TATATCAATAAGAAAACCTTTTATAGTAATGAATAAAGGAAAGGCAATT					
							Section 17
	(769)	769	780	790	800		816
ClareAJ251507	(609)	TTCCGATTTCAGTGCCACCTCTGCCTTGTATATTTTAACTCCACTAAAC					
huNall18 (AK)	(305)	TTCCGATTTCAGTGCCACCTCTGCCTTGTATATTTTAACTCCACTAAAC					
JeongAF225987	(769)	TTCCGATTTCAGTGCCACCTCTGCCTTGTATATTTTAACTCCACTAAAC					
Consensus	(769)	TTCCGATTTCAGTGCCACCTCTGCCTTGTATATTTTAACTCCACTAAAC					
							Section 18
	(817)	817	830	840	850		864
ClareAJ251507	(657)	CCTGTTAGGAAAATTGCTATCAAGATTTTGGTACATTCTTTATTTCAGC					
huNall18 (AK)	(353)	CCTGTTAGGAAAATTGCTATCAAGATTTTGGTACATTCTTTATTTCAGC					
JeongAF225987	(817)	CCTGTTAGGAAAATTGCTATCAAGATTTTGGTACATTCTTTATTTCAGC					
Consensus	(817)	CCTGTTAGGAAAATTGCTATCAAGATTTTGGTACATTCTTTATTTCAGC					
							Section 19
	(865)	865	870	880	890	900	912
ClareAJ251507	(705)	ATGCTTATCATGTGCACTATTTTGACCAACTGTGTATTTATGACCTTG					
huNall18 (AK)	(401)	ATGCTTATCATGTGCACTATTTTGACCAACTGTGTATTTATGACCTTG					
JeongAF225987	(865)	ATGCTTATCATGTGCACTATTTTGACCAACTGTGTATTTATGACCTTG					
Consensus	(865)	ATGCTTATCATGTGCACTATTTTGACCAACTGTGTATTTATGACCTTG					
							Section 20
	(913)	913	920	930	940	950	960
ClareAJ251507	(753)	AGCAACCCTCCTGACTGGACAAAGAATGTAGAGTACACATTCCTGGA					
huNall18 (AK)	(449)	AGCAACCCTCCTGACTGGACAAAGAATGTAGAGTACACATTCCTGGA					
JeongAF225987	(913)	AGCAACCCTCCTGACTGGACAAAGAATGTAGAGTACACATTCCTGGA					
Consensus	(913)	AGCAACCCTCCTGACTGGACAAAGAATGTAGAGTACACATTCCTGGA					
							Section 21
	(961)	961	970	980	990		1008
ClareAJ251507	(801)	ATCTATACCTTTGAGTCACCTTATAAAAAATCTTGGCAAGAGGGTTTTGC					
huNall18 (AK)	(497)	ATCTATACCTTTGAGTCACCTTATAAAAAATCTTGGCAAGAGGGTTTTGC					
JeongAF225987	(961)	ATCTATACCTTTGAGTCACCTTATAAAAAATCTTGGCAAGAGGGTTTTGC					
Consensus	(961)	ATCTATACCTTTGAGTCACCTTATAAAAAATCTTGGCAAGAGGGTTTTGC					

Section 22

	(1009)	1009	1020	1030	1040	1056
ClareAJ251507	(849)	TTAGAAGATTTTACGTTTCTTCGTGATCCATGGAAC	TGGCTGGATTTC			
huNall18 (AK)	(545)	TTAGAAGATTTTACGTTTCTTCGTGATCCATGGAAC	TGGCTGGATTTC			
JeongAF225987	(1009)	TTAGAAGATTTTACGTTTCTTCGTGATCCATGGAAC	TGGCTGGATTTC			
Consensus	(1009)	TTAGAAGATTTTACGTTTCTTCGTGATCCATGGAAC	TGGCTGGATTTC			

Section 23

	(1057)	1057	1070	1080	1090	1104
ClareAJ251507	(897)	AGTGTCAATTGTGATGGCGTATGTACAGAAATTTGTAAAGCCTAGGCAAT				
huNall18 (AK)	(593)	AGTGTCAATTGTGATGGCGTATGTACAGAAATTTGTAAAGCCTAGGCAAT				
JeongAF225987	(1057)	AGTGTCAATTGTGATGGCGTATGTACAGAGTTTGTGGACCTGGGCAAT				
Consensus	(1057)	AGTGTCAATTGTGATGGCGTATGTAACAGAAATTTGTAAAGCCTAGGCAAT				

Section 24

	(1105)	1105	1110	1120	1130	1140	1152
ClareAJ251507	(945)	GTCTCAGCCCTTCGAACATTCAGAGTCTTGAGAGCCTTGAAACATT					
huNall18 (AK)	(641)	GTCTCAGCCCTTCGAACATTCAGAGTCTTGAGAGCCTTGAAACATT					
JeongAF225987	(1105)	GTCTCAGCGTTGAGAACATTCAGAGTCTCCGAGCACTGAAACAATT					
Consensus	(1105)	GTTTCAGCCCTTCGAACATTCAGAGTCTTGAGAGCCTTGAAACATT					

Section 25

	(1153)	1153	1160	1170	1180	1190	1200
ClareAJ251507	(993)	TCCTGTATTCCAGGTTTAAAGACCATTGTGGGGGCCCTGATCCAGTCG					
huNall18 (AK)	(689)	TCCTGTATTCCAGGTTTAAAGACCATTGTGGGGGCCCTGATCCAGTCG					
JeongAF225987	(1153)	TCAGTCATTCCAGGTTTAAAGACCATTGTGGGGGCCCTGATCCAGTCG					
Consensus	(1153)	TCTGTAATTCCAGGTTTAAAGACCATTGTGGGGGCCCTGATCCAGTCG					

Section 26

	(1201)	1201	1210	1220	1230	1248
ClareAJ251507	(1041)	GTAAAGAAGCTTTCTGATGTGATGATCCTGACTGTGTTCTGTCTGAGC				
huNall18 (AK)	(737)	GTAAAGAAGCTTTCTGATGTGATGATCCTGACTGTGTTCTGTCTGAGC				
JeongAF225987	(1201)	GTAAAGAAGCTTTCTGATGTGATGATCCTGACTGTGTTCTGTCTGAGC				
Consensus	(1201)	GTAAAGAAGCTTTCTGATGTGATGATCCTGACTGTGTTCTGTCTGAGC				

Section 27

	(1249)	1249	1260	1270	1280	1296
ClareAJ251507	(1089)	GTGTTTGCTCTCATTGGGCTGCAGCTGTTTCATGGGCAATCTGAGGAAT				
huNall18 (AK)	(785)	GTGTTTGCTCTCATTGGGCTGCAGCTGTTTCATGGGCAATCTGAGGAAT				
JeongAF225987	(1249)	GTGTTTGCTCTCATTGGGCTGCAGCTGTTTCATGGGCAATCTGAGGAAT				
Consensus	(1249)	GTGTTTGCTCTCATTGGGCTGCAGCTGTTTCATGGGCAATCTGAGGAAT				

Section 28

	(1297)	1297	1310	1320	1330	1344
ClareAJ251507	(1137)	AAATGTTTGTCAGTGGCCCCCAAGCGATTCTGCTTTTGAACCAACACC				
huNall18 (AK)	(833)	AAATGTTTGTCAGTGGCCCCCAAGCGATTCTGCTTTTGAACCAACACC				
JeongAF225987	(1297)	AAATGTTTGTCAGTGGCCCCCAAGCGATTCTGCTTTTGAACCAACACC				
Consensus	(1297)	AAATGTTTGTCAGTGGCCCCCAAGCGATTCTGCTTTTGAACCAACACC				

Section 29

	(1345)	1345	1350	1360	1370	1380	1392
ClareAJ251507 (1185)		ACTTCCTACTTTAATGGCACAATGGATTCAAATGGGACATTTGTTAAT					
huNall18 (AK) (881)		ACTTCCTACTTTAATGGCACAATGGATTCAAATGGGACATTTGTTAAT					
JeongAF225987 (1345)		ACTTCCTACTTTAATGGCACAATGGATTCAAATGGGACATTTGTTAAT					
Consensus (1345)		ACTTCCTACTTTAATGGCACAATGGATTCAAATGGGACATTTGTTAAT					

Section 30

	(1393)	1393	1400	1410	1420	1430	1440
ClareAJ251507 (1233)		GTAACAATGAGCACATTTAACTGGAAGGATTACATTGGAGATGACAGT					
huNall18 (AK) (929)		GTAACAATGAGCACATTTAACTGGAAGGATTACATTGGAGATGACAGT					
JeongAF225987 (1393)		GTAACAATGAGCACATTTAACTGGAAGGATTACATTGGAGATGACAGT					
Consensus (1393)		GTAACAATGAGCACATTTAACTGGAAGGATTACATTGGAGATGACAGT					

Section 31

	(1441)	1441	1450	1460	1470	1488
ClareAJ251507 (1281)		CACTTTTATGTTTTGGATGGGCAAAAAGACCCTTTACTCTGTGGAAAT				
huNall18 (AK) (977)		CACTTTTATGTTTTGGATGGGCAAAAAGACCCTTTACTCTGTGGAAAT				
JeongAF225987 (1441)		CACTTTTATGTTTTGGATGGGCAAAAAGACCCTTTACTCTGTGGAAAT				
Consensus (1441)		CACTTTTATGTTTTGGATGGGCAAAAAGACCCTTTACTCTGTGGAAAT				

Section 32

	(1489)	1489	1500	1510	1520	1536
ClareAJ251507 (1329)		GGCTCAGATGCAGGCCAGTGTCCAGAAGGATACATCTGTGTGAAGGCT				
huNall18 (AK) (1025)		GGCTCAGATGCAGGCCAGTGTCCAGAAGGATACATCTGTGTGAAGGCT				
JeongAF225987 (1489)		GGCTCAGATGCAGGCCAGTGTCCAGAAGGATACATCTGTGTGAAGGCT				
Consensus (1489)		GGCTCAGATGCAGGCCAGTGTCCAGAAGGATACATCTGTGTGAAGGCT				

Section 33

	(1537)	1537	1550	1560	1570	1584
ClareAJ251507 (1377)		GGTCGAAACCCCAACTATGGCTACACAAGCTTTGACACCTTTAGCTGG				
huNall18 (AK) (1073)		GGTCGAAACCCCAACTATGGCTACACAAGCTTTGACACCTTTAGCTGG				
JeongAF225987 (1537)		GGTCGAAACCCCAACTATGGCTACACAAGCTTTGACACCTTTAGCTGG				
Consensus (1537)		GGTCGAAACCCCAACTATGGCTACACAAGCTTTGACACCTTTAGCTGG				

Section 34

	(1585)	1585	1590	1600	1610	1620	1632
ClareAJ251507 (1425)		GCTTTCCTGTCTCTATTTTCGACTCATGACTCAAGATTATGGGAAAAT					
huNall18 (AK) (1121)		GCTTTCCTGTCTCTATTTTCGACTCATGACTCAAGATTATGGGAAAAT					
JeongAF225987 (1585)		GCTTTCCTGTCTCTATTTTCGACTCATGACTCAAGATTATGGGAAAAT					
Consensus (1585)		GCTTTCCTGTCTCTATTTTCGACTCATGACTCAAGACTACTGGGAAAAT					

Section 35

	(1633)	1633	1640	1650	1660	1670	1680
ClareAJ251507 (1473)		CTTTACCAGTTGACATTACGTGCTGCTGGGAAAACATACATGATATTT					
huNall18 (AK) (1169)		CTTTACCAGTTGACATTACGTGCTGCTGGGAAAACATACATGATATTT					
JeongAF225987 (1633)		CTTTACCAGTTGACATTACGTGCTGCTGGGAAAACATACATGATATTT					
Consensus (1633)		CTTTACCAGTTGACATTACGTGCTGCTGGGAAAACATACATGATATTT					

Section 36							
	(1681)	1681	1690	1700	1710	1728	
ClareAJ251507	(1521)	TTTGTCTCTGGTCATTTTCTTGGGCTCATTTTATTTGGTGAATTTGATC					
huNall18 (AK)	(1217)	TTTGTCTCTGGTCATTTTCTTGGGCTCATTTTATTTGGTGAATTTGATC					
JeongAF225987	(1681)	TTTGTCTCTGGTCATTTTCTTGGGCTCATTTTATTTGGTGAATTTGATC					
Consensus	(1681)	TTTGTCTCTGGTCATTTTCTTGGGCTCATTTTATTTGGTGAATTTGATC					
Section 37							
	(1729)	1729	1740	1750	1760	1776	
ClareAJ251507	(1569)	CTGGCTGTGGTGGCCATGGCCTATGAGGAGCAGAATCAGGCCACCTTG					
huNall18 (AK)	(1265)	CTGGCTGTGGTGGCCATGGCCTATGAGGAGCAGAATCAGGCCACCTTG					
JeongAF225987	(1729)	CTGGCTGTGGTGGCCATGGCCTATGAGGAGCAGAATCAGGCCACCTTG					
Consensus	(1729)	CTGGCTGTGGTGGCCATGGCCTATGAGGAGCAGAATCAGGCCACCTTG					
Section 38							
	(1777)	1777	1790	1800	1810	1824	
ClareAJ251507	(1617)	GAAGAAGCAGAACAAAAAGAGGCCGAATTTTCAGCAGATGCTCGAACAG					
huNall18 (AK)	(1313)	GAAGAAGCAGAACAAAAAGAGGCCGAATTTTCAGCAGATGCTCGAACAG					
JeongAF225987	(1777)	GAAGAAGCAGAACAAAAAGAGGCCGAATTTTCAGCAGATGCTCGAACAG					
Consensus	(1777)	GAAGAAGCAGAACAAAAAGAGGCCGAATTTTCAGCAGATGCTCGAACAG					
Section 39							
	(1825)	1825	1830	1840	1850	1860	1872
ClareAJ251507	(1665)	CTTAAAAAGCAACAGGAAGAAGCTCAGGCAGTTGCGGCAGCATCAGCT					
huNall18 (AK)	(1361)	CTTAAAAAGCAACAGGAAGAAGCTCAGGCAGTTGCGGCAGCATCAGCT					
JeongAF225987	(1825)	CTTAAAAAGCAACAGGAAGAAGCTCAGGCAGTTGCGGCAGCATCAGCT					
Consensus	(1825)	CTTAAAAAGCAACAGGAAGAAGCTCAGGCAGTTGCGGCAGCATCAGCT					
Section 40							
	(1873)	1873	1880	1890	1900	1910	1920
ClareAJ251507	(1713)	GCTTCAAGAGATTTTCAGTGGAGTAGGTGGGTTAGGAGAGCTGTTGGAA					
huNall18 (AK)	(1409)	GCTTCAAGAGATTTTCAGTGGAGTAGGTGGGTTAGGAGAGCTGTTGGAA					
JeongAF225987	(1873)	GCTTCAAGAGATTTTCAGTGGAGTAGGTGGGTTAGGAGAGCTGTTGGAA					
Consensus	(1873)	GCTTCAAGAGATTTTCAGTGGAGTAGGTGGGTTAGGAGAGCTGTTGGAA					
Section 41							
	(1921)	1921	1930	1940	1950	1968	
ClareAJ251507	(1761)	AGTTCTTCAGAAGCATCAAAGTTGAGTTCCAAAAGTGCTAAAGAATGG					
huNall18 (AK)	(1457)	AGTTCTTCAGAAGCATCAAAGTTGAGTTCCAAAAGTGCTAAAGAATGG					
JeongAF225987	(1921)	AGTTCTTCAGAAGCATCAAAGTTGAGTTCCAAAAGTGCTAAAGAATGG					
Consensus	(1921)	AGTTCTTCAGAAGCATCAAAGTTGAGTTCCAAAAGTGCTAAAGAATGG					
Section 42							
	(1969)	1969	1980	1990	2000	2016	
ClareAJ251507	(1809)	AGGAACCGAGGAAGAAAAGAAGACAGAGAGAGCACCTTGAAGGAAAC					
huNall18 (AK)	(1505)	AGGAACCGAGGAAGAAAAGAAGACAGAGAGAGCACCTTGAAGGAAAC					
JeongAF225987	(1969)	AGGAACCGAGGAAGAAAAGAAGACAGAGAGAGCACCTTGAAGGAAAC					
Consensus	(1969)	AGGAACCGAGGAAGAAAAGAAGACAGAGAGAGCACCTTGAAGGAAAC					

Section 43

	(2017)	2017	2030	2040	2050	2064
ClareAJ251507 (1857)		AACAAAGGAGAGAGAGACAGCTTTCCCAAATCCGAATCTGAAGACAGC				
huNall18 (AK) (1553)		AACAAAGGAGAGAGAGAGACAGCTTTCCCAAATCCGAATCTGAAGACAGC				
JeongAF225987 (2017)		AACAAAGGAGAGAGAGAGACAGCTTTCCCAAATCCGAATCTGAAGACAGC				
Consensus (2017)		AACAAAGGAGAGAGAGAGACAGCTTTCCCAAATCCGAATCTGAAGACAGC				

Section 44

	(2065)	2065	2070	2080	2090	2100	2112
ClareAJ251507 (1905)		GTCAAAAGAAGCAGCTTCCTTTTCTCCATGGATGGAAACAGACTGACC					
huNall18 (AK) (1601)		GTCAAAAGAAGCAGCTTCCTTTTCTCCATGGATGGAAACAGACTGACC					
JeongAF225987 (2065)		GTCAAAAGAAGCAGCTTCCTTTTCTCCATGGATGGAAACAGACTGACC					
Consensus (2065)		GTCAAAAGAAGCAGCTTCCTTTTCTCCATGGATGGAAACAGACTGACC					

Section 45

	(2113)	2113	2120	2130	2140	2150	2160
ClareAJ251507 (1953)		AGTGACAAAAAATTCTGCTCCCCTCATCAGTCTCTCTTGAGTATCCGT					
huNall18 (AK) (1649)		AGTGACAAAAAATTCTGCTCCCCTCATCAGTCTCTCTTGAGTATCCGT					
JeongAF225987 (2113)		AGTGACAAAAAATTCTGCTCCCCTCATCAGTCTCTCTTGAGTATCCGT					
Consensus (2113)		AGTGACAAAAAATTCTGCTCCCCTCATCAGTCTCTCTTGAGTATCCGT					

Section 46

	(2161)	2161	2170	2180	2190	2208
ClareAJ251507 (2001)		GGCTCCCTGTTTTCCCAAGACGCAATAGCAAAACAAGCATTTTCAGT				
huNall18 (AK) (1697)		GGCTCCCTGTTTTCCCAAGACGCAATAGCAAAACAAGCATTTTCAGT				
JeongAF225987 (2161)		GGCTCCCTGTTTTCCCAAGACGCAATAGCAAAACAAGCATTTTCAGT				
Consensus (2161)		GGCTCCCTGTTTTCCCAAGACGCAATAGCAAAACAAGCATTTTCAGT				

Section 47

	(2209)	2209	2220	2230	2240	2256
ClareAJ251507 (2049)		TTCAGAGGTCGGGCAAAGGATGTTGGATCTGAAAATGACTTTGCTGAT				
huNall18 (AK) (1745)		TTCAGAGGTCGGGCAAAGGATGTTGGATCTGAAAATGACTTTGCTGAT				
JeongAF225987 (2209)		TTCAGAGGTCGGGCAAAGGATGTTGGATCTGAAAATGACTTTGCTGAT				
Consensus (2209)		TTCAGAGGTCGGGCAAAGGATGTTGGATCTGAAAATGACTTTGCTGAT				

Section 48

	(2257)	2257	2270	2280	2290	2304
ClareAJ251507 (2097)		GATGAACACAGCACATTTGAAGACAGCGAAAGCAGGAGAGACTCACTG				
huNall18 (AK) (1793)		GATGAACACAGCACATTTGAAGACAGCGAAAGCAGGAGAGACTCACTG				
JeongAF225987 (2257)		GATGAACACAGCACATTTGAAGACAGCGAAAGCAGGAGAGACTCACTG				
Consensus (2257)		GATGAACACAGCACATTTGAAGACAGCGAAAGCAGGAGAGACTCACTG				

Section 49

	(2305)	2305	2310	2320	2330	2340	2352
ClareAJ251507 (2145)		TTTGTGCCGCACAGACATGGAGAGCGACGCAACAGTAACG-----					
huNall18 (AK) (1841)		TTTGTGCCGCACAGACATGGAGAGCGACGCAACAGTAACGTTAGTCAG					
JeongAF225987 (2305)		TTTGTGCCGCACAGACATGGAGAGCGACGCAACAGTAACGTTAGTCAG					
Consensus (2305)		TTTGTGCCGCACAGACATGGAGAGCGACGCAACAGTAACGTTAGTCAG					

						Section 50
	(2353)	2353	2360	2370	2380	2390 2400
ClareAJ251507 (2185)		-----	-----	-----	-----	-----
huNall18 (AK) (1889)		GCCAGTATGTCATCCAGGATGGTGCCAGGGCTTCCAGCAAATGGGAAG				
JeongAF225987 (2353)		GCCAGTATGTCATCCAGGATGGTGCCAGGGCTTCCAGCAAATGGGAAG				
Consensus (2353)		GCCAGTATGTCATCCAGGATGGTGCCAGGGCTTCCAGCAAATGGGAAG				
						Section 51
	(2401)	2401	2410	2420	2430	2448
ClareAJ251507 (2185)		-----	-----	-----	-----	-----
huNall18 (AK) (1937)		ATGCACAGCACTGTGGATTGCAATGGTGTGGTTTCCTTGGTGGGTGGA				
JeongAF225987 (2401)		ATGCACAGCACTGTGGATTGCAATGGTGTGGTTTCCTTGGTGGGTGGA				
Consensus (2401)		ATGCACAGCACTGTGGATTGCAATGGTGTGGTTTCCTTGGTGGGTGGA				
						Section 52
	(2449)	2449	2460	2470	2480	2496
ClareAJ251507 (2185)		-----	-----	-----	-----	-----GCACC
huNall18 (AK) (1985)		CCTTCAGCTCTAACGTCACCTACTGGACAACCTTCCCCCAGAGGGGCACC				
JeongAF225987 (2449)		CCTTCAGCTCTAACGTCACCTACTGGACAACCTTCCCCCAGAGGGGCACC				
Consensus (2449)		CCTTCAGCTCTAACGTCACCTACTGGACAACCTTCCCCCAGAGGGGCACC				
						Section 53
	(2497)	2497	2510	2520	2530	2544
ClareAJ251507 (2190)		ACCACGAAACGGAAGTCAGAAAGAGAAGGTTAAGCTCTTACCAGATT				
huNall18 (AK) (2033)		ACCACGAAACGGAAGTCAGAAAGAGAAGGTTAAGCTCTTACCAGATT				
JeongAF225987 (2497)		ACCACGAAACGGAAGTCAGAAAGAGAAGGTTAAGCTCTTACCAGATT				
Consensus (2497)		ACCACGAAACGGAAGTCAGAAAGAGAAGGTTAAGCTCTTACCAGATT				
						Section 54
	(2545)	2545	2550	2560	2570	2580 2592
ClareAJ251507 (2238)		TCAATGGAGATGCTGGAGGATTCTCTGGAAGGCAAAGAGCCGTGAGC				
huNall18 (AK) (2081)		TCAATGGAGATGCTGGAGGATTCTCTGGAAGGCAAAGAGCCGTGAGC				
JeongAF225987 (2545)		TCAATGGAGATGCTGGAGGATTCTCTGGAAGGCAAAGAGCCGTGAGC				
Consensus (2545)		TCAATGGAGATGCTGGAGGATTCTCTGGAAGGCAAAGAGCCGTGAGC				
						Section 55
	(2593)	2593	2600	2610	2620	2630 2640
ClareAJ251507 (2286)		ATAGCCAGCATTTCTGACCAACACAATGGAAGAACTTGAAGAATCTAGA				
huNall18 (AK) (2129)		ATAGCCAGCATTTCTGACCAACACAATGGAAGAACTTGAAGAATCTAGA				
JeongAF225987 (2593)		ATAGCCAGCATTTCTGACCAACACAATGGAAGAACTTGAAGAATCTAGA				
Consensus (2593)		ATAGCCAGCATTTCTGACCAACACAATGGAAGAACTTGAAGAATCTAGA				
						Section 56
	(2641)	2641	2650	2660	2670	2688
ClareAJ251507 (2334)		CAGAAATGTCCGCCATGCTGGTATAGATTTGCCAATGTGTTCTTGATC				
huNall18 (AK) (2177)		CAGAAATGTCCGCCATGCTGGTATAGATTTGCCAATGTGTTCTTGATC				
JeongAF225987 (2641)		CAGAAATGTCCGCCATGCTGGTATAGATTTGCCAATGTGTTCTTGATC				
Consensus (2641)		CAGAAATGTCCGCCATGCTGGTATAGATTTGCCAATGTGTTCTTGATC				

Section 57

	(2689)	2689	2700	2710	2720	2736
ClareAJ251507 (2382)		TGGGACTGCTGTGATGCATGGTTAAAAGTAAAACATCTTGTGAATTTA				
huNall18 (AK) (2225)		TGGGACTGCTGTGATGCATGGTTAAAAGTAAAACATCTTGTGAATTTA				
JeongAF225987 (2689)		TGGGACTGCTGTGATGCATGGTTAAAAGTAAAACATCTTGTGAATTTA				
Consensus (2689)		TGGGACTGCTGTGATGCATGGTTAAAAGTAAAACATCTTGTGAATTTA				

Section 58

	(2737)	2737	2750	2760	2770	2784
ClareAJ251507 (2430)		ATTGTTATGGATCCATTTGTTGATCTTGCCATCACTATTTGCATTGTC				
huNall18 (AK) (2273)		ATTGTTATGGATCCATTTGTTGATCTTGCCATCACTATTTGCATTGTC				
JeongAF225987 (2737)		ATTGTTATGGATCCATTTGTTGATCTTGCCATCACTATTTGCATTGTC				
Consensus (2737)		ATTGTTATGGATCCATTTGTTGATCTTGCCATCACTATTTGCATTGTC				

Section 59

	(2785)	2785	2790	2800	2810	2820	2832
ClareAJ251507 (2478)		TTAAATACCCCTCTTTATGGCCATGGAGCACTACCCCATGACTGAGCAA					
huNall18 (AK) (2321)		TTAAATACCCCTCTTTATGGCCATGGAGCACTACCCCATGACTGAGCAA					
JeongAF225987 (2785)		TTAAATACCCCTCTTTATGGCCATGGAGCACTACCCCATGACTGAGCAA					
Consensus (2785)		TTAAATACCCCTCTTTATGGCCATGGAGCACTACCCCATGACTGAGCAA					

Section 60

	(2833)	2833	2840	2850	2860	2870	2880
ClareAJ251507 (2526)		TTCAGTAGTGTGTTGACTGTAGGAAACCTGGTCTTTACTGGGATTTTC					
huNall18 (AK) (2369)		TTCAGTAGTGTGTTGACTGTAGGAAACCTGGTCTTTACTGGGATTTTC					
JeongAF225987 (2833)		TTCAGTAGTGTGTTGACTGTAGGAAACCTGGTCTTTACTGGGATTTTC					
Consensus (2833)		TTCAGTAGTGTGTTGACTGTAGGAAACCTGGTCTTTACTGGGATTTTC					

Section 61

	(2881)	2881	2890	2900	2910	2928
ClareAJ251507 (2574)		ACAGCAGAAATGGTTCTCAAGATCATTGCCATGGATCCTTATTACTAT				
huNall18 (AK) (2417)		ACAGCAGAAATGGTTCTCAAGATCATTGCCATGGATCCTTATTACTAT				
JeongAF225987 (2881)		ACAGCAGAAATGGTTCTCAAGATCATTGCCATGGATCCTTATTACTAT				
Consensus (2881)		ACAGCAGAAATGGTTCTCAAGATCATTGCCATGGATCCTTATTACTAT				

Section 62

	(2929)	2929	2940	2950	2960	2976
ClareAJ251507 (2622)		TTCCAAGAAGGCTGGAATATCTTTGATGGAATTATTGTCAGCCTCAGT				
huNall18 (AK) (2465)		TTCCAAGAAGGCTGGAATATCTTTGATGGAATTATTGTCAGCCTCAGT				
JeongAF225987 (2929)		TTCCAAGAAGGCTGGAATATCTTTGATGGAATTATTGTCAGCCTCAGT				
Consensus (2929)		TTCCAAGAAGGCTGGAATATCTTTGATGGAATTATTGTCAGCCTCAGT				

Section 63

	(2977)	2977	2990	3000	3010	3024
ClareAJ251507 (2670)		TTAATGGAGCTTGGTCTGTCAAATGTGGAGGGATTGTCTGTACTGCGA				
huNall18 (AK) (2513)		TTAATGGAGCTTGGTCTGTCAAATGTGGAGGGATTGTCTGTACTGCGA				
JeongAF225987 (2977)		TTAATGGAGCTTGGTCTGTCAAATGTGGAGGGATTGTCTGTACTGCGA				
Consensus (2977)		TTAATGGAGCTTGGTCTGTCAAATGTGGAGGGATTGTCTGTACTGCGA				

Section 64							
	(3025)	3025	3030	3040	3050	3060	3072
ClareAJ251507	(2718)	TCATTCAGACTGCTTAGAGTTTTCAAGTTGGCAAATCCTGGCCCCACA					
huNall18 (AK)	(2561)	TCATTCAGACTGCTTAGAGTTTTCAAGTTGGCAAATCCTGGCCCCACA					
JeongAF225987	(3025)	TCATTCAGACTGCTTAGAGTTTTCAAGTTGGCAAATCCTGGCCCCACA					
Consensus	(3025)	TCATTCAGACTGCTTAGAGTTTTCAAGTTGGCAAATCCTGGCCCCACA					
Section 65							
	(3073)	3073	3080	3090	3100	3110	3120
ClareAJ251507	(2766)	CTAAATATGCTAATTAAGATCATTTGGCAATTCTGTGGGGGCTCTAGGA					
huNall18 (AK)	(2609)	CTAAATATGCTAATTAAGATCATTTGGCAATTCTGTGGGGGCTCTAGGA					
JeongAF225987	(3073)	CTAAATATGCTAATTAAGATCATTTGGCAATTCTGTGGGGGCTCTAGGA					
Consensus	(3073)	CTAAATATGCTAATTAAGATCATTTGGCAATTCTGTGGGGGCTCTAGGA					
Section 66							
	(3121)	3121	3130	3140	3150		3168
ClareAJ251507	(2814)	AACCTCACCTTGGTGTGGCCATCATCGTCTTCATTTTTGCTGTGGTC					
huNall18 (AK)	(2657)	AACCTCACCTTGGTGTGGCCATCATCGTCTTCATTTTTGCTGTGGTC					
JeongAF225987	(3121)	AACCTCACCTTGGTGTGGCCATCATCGTCTTCATTTTTGCTGTGGTC					
Consensus	(3121)	AACCTCACCTTGGTGTGGCCATCATCGTCTTCATTTTTGCTGTGGTC					
Section 67							
	(3169)	3169	3180	3190	3200		3216
ClareAJ251507	(2862)	GGCATGCAGCTCTTTGGTAAGAGCTACAAAGAATGTGTCTGCAAGATC					
huNall18 (AK)	(2705)	GGCATGCAGCTCTTTGGTAAGAGCTACAAAGAATGTGTCTGCAAGATC					
JeongAF225987	(3169)	GGCATGCAGCTCTTTGGTAAGAGCTACAAAGAATGTGTCTGCAAGATC					
Consensus	(3169)	GGCATGCAGCTCTTTGGTAAGAGCTACAAAGAATGTGTCTGCAAGATC					
Section 68							
	(3217)	3217	3230	3240	3250		3264
ClareAJ251507	(2910)	AATGATGACTGTACGCTCCCACGGTGGCACATGAACGACTTCTTCCAC					
huNall18 (AK)	(2753)	AATGATGACTGTACGCTCCCACGGTGGCACATGAACGACTTCTTCCAC					
JeongAF225987	(3217)	AATGATGACTGTACGCTCCCACGGTGGCACATGAACGACTTCTTCCAC					
Consensus	(3217)	AATGATGACTGTACGCTCCCACGGTGGCACATGAACGACTTCTTCCAC					
Section 69							
	(3265)	3265	3270	3280	3290	3300	3312
ClareAJ251507	(2958)	TCCTTCCTGATTGTGTTCCGCGTGCTGTGTGGAGAGTGGATAGAGACC					
huNall18 (AK)	(2801)	TCCTTCCTGATTGTGTTCCGCGTGCTGTGTGGAGAGTGGATAGAGACC					
JeongAF225987	(3265)	TCCTTCCTGATTGTGTTCCGCGTGCTGTGTGGAGAGTGGATAGAGACC					
Consensus	(3265)	TCCTTCCTGATTGTGTTCCGCGTGCTGTGTGGAGAGTGGATAGAGACC					
Section 70							
	(3313)	3313	3320	3330	3340	3350	3360
ClareAJ251507	(3006)	ATGTGGGACTGTATGGAGGTCGCTGGCCAAACCATGTGCCTTATTGTT					
huNall18 (AK)	(2849)	ATGTGGGACTGTATGGAGGTCGCTGGCCAAACCATGTGCCTTATTGTT					
JeongAF225987	(3313)	ATGTGGGACTGTATGGAGGTCGCTGGCCAAACCATGTGCCTTATTGTT					
Consensus	(3313)	ATGTGGGACTGTATGGAGGTCGCTGGCCAAACCATGTGCCTTATTGTT					

Section 71							
	(3361)	3361	3370	3380	3390	3408	
ClareAJ251507	(3054)	TTCATGTTGGTCATGGTCATTGGAAACCTTGTGGTTCTGAACCTCTTT					
huNall118 (AK)	(2897)	TTCATGTTGGTCATGGTCATTGGAAACCTTGTGGTTCTGAACCTCTTT					
JeongAF225987	(3361)	TTCATGTTGGTCATGGTCATTGGAAACCTTGTGGTTCTGAACCTCTTT					
Consensus	(3361)	TTCATGTTGGTCATGGTCATTGGAAACCTTGTGGTTCTGAACCTCTTT					
Section 72							
	(3409)	3409	3420	3430	3440	3456	
ClareAJ251507	(3102)	CTGGCCTTATTGTTGAGTTCATTTAGCTCAGACAACCTTGCTGCTACT					
huNall118 (AK)	(2945)	CTGGCCTTATTGTTGAGTTCATTTAGCTCAGACAACCTTGCTGCTACT					
JeongAF225987	(3409)	CTGGCCTTATTATTGAGTTCATTTAGCTCAGACAACCTTGCTGCTACT					
Consensus	(3409)	CTGGCCTTATTGTTGAGTTCATTTAGCTCAGACAACCTTGCTGCTACT					
Section 73							
	(3457)	3457	3470	3480	3490	3504	
ClareAJ251507	(3150)	GATGATGACAATGAAATGAATAATCTGCAGATTGCAGTAGGAAGAATG					
huNall118 (AK)	(2993)	GATGATGACAATGAAATGAATAATCTGCAGATTGCAGTAGGAAGAATG					
JeongAF225987	(3457)	GATGATGACAATGAAATGAATAATCTGCAGATTGCAGTAGGAAGAATG					
Consensus	(3457)	GATGATGACAATGAAATGAATAATCTGCAGATTGCAGTAGGAAGAATG					
Section 74							
	(3505)	3505	3510	3520	3530	3540	3552
ClareAJ251507	(3198)	CAAAAGGGAATTGATTATGTGAAAAATAAGATGCGGGAGTGTTTCCAA					
huNall118 (AK)	(3041)	CAAAAGGGAATTGATTATGTGAAAAATAAGATGCGGGAGTGTTTCCAA					
JeongAF225987	(3505)	CAAAAGGGAATTGATTATGTGAAAAATAAGATGCGGGAGTGTTTCCAA					
Consensus	(3505)	CAAAAGGGAATTGATTATGTGAAAAATAAGATGCGGGAGTGTTTCCAA					
Section 75							
	(3553)	3553	3560	3570	3580	3590	3600
ClareAJ251507	(3246)	AAAGCCTTTTTTAGAAAGCCAAAAGTTATAGAAATCCATGAAGGCAAT					
huNall118 (AK)	(3089)	AAAGCCTTTTTTAGAAAGCCAAAAGTTATAGAAATCCATGAAGGCAAT					
JeongAF225987	(3553)	AAAGCCTTTTTTAGAAAGCCAAAAGTTATAGAAATCCATGAAGGCAAT					
Consensus	(3553)	AAAGCCTTTTTTAGAAAGCCAAAAGTTATAGAAATCCATGAAGGCAAT					
Section 76							
	(3601)	3601	3610	3620	3630	3648	
ClareAJ251507	(3294)	AAGATAGACAGCTGCATGTCCAATAATACTGGAATTGAAATAAGCAAA					
huNall118 (AK)	(3137)	AAGATAGACAGCTGCATGTCCAATAATACTGGAATTGAAATAAGCAAA					
JeongAF225987	(3601)	AAGATAGACAGCTGCATGTCCAATAATACTGGAATTGAAATAAGCAAA					
Consensus	(3601)	AAGATAGACAGCTGCATGTCCAATAATACTGGAATTGAAATAAGCAAA					
Section 77							
	(3649)	3649	3660	3670	3680	3696	
ClareAJ251507	(3342)	GAGCTTAATTATCTTAGAGATGGGAATGGAACCACCAGTGGTGTAGGT					
huNall118 (AK)	(3185)	GAGCTTAATTATCTTAGAGATGGGAATGGAACCACCAGTGGTGTAGGT					
JeongAF225987	(3649)	GAGCTTAATTATCTTAGAGATGGGAATGGAACCACCAGTGGTGTAGGT					
Consensus	(3649)	GAGCTTAATTATCTTAGAGATGGGAATGGAACCACCAGTGGTGTAGGT					

Section 78

	(3697)	3697	3710	3720	3730	3744
ClareAJ251507	(3390)	ACTGGAAGCAGTGTTGAAAAATACGTAATCGATGAAAATGATTATATG				
huNall18 (AK)	(3233)	ACTGGAAGCAGTGTTGAAAAATACGTAATCGATGAAAATGATTATATG				
JeongAF225987	(3697)	ACTGGAAGCAGTGTTGAAAAATACGTAATCGATGAAAATGATTATATG				
Consensus	(3697)	ACTGGAAGCAGTGTTGAAAAATACGTAATCGATGAAAATGATTATATG				

Section 79

	(3745)	3745	3750	3760	3770	3780	3792
ClareAJ251507	(3438)	TCATTCATAAACAACCCAGCCTCACCGTCACAGTGCCAATTGCTGTT					
huNall18 (AK)	(3281)	TCATTCATAAACAACCCAGCCTCACCGTCACAGTGCCAATTGCTGTT					
JeongAF225987	(3745)	TCATTCATAAACAACCCAGCCTCACCGTCACAGTGCCAATTGCTGTT					
Consensus	(3745)	TCATTCATAAACAACCCAGCCTCACCGTCACAGTGCCAATTGCTGTT					

Section 80

	(3793)	3793	3800	3810	3820	3830	3840
ClareAJ251507	(3486)	GGAGAGTCTGACTTTGAAAACCTTAAATACTGAAGAGTTCAGCAGTGAG					
huNall18 (AK)	(3329)	GGAGAGTCTGACTTTGAAAACCTTAAATACTGAAGAGTTCAGCAGTGAG					
JeongAF225987	(3793)	GGAGAGTCTGACTTTGAAAACCTTAAATACTGAAGAGTTCAGCAGTGAG					
Consensus	(3793)	GGAGAGTCTGACTTTGAAAACCTTAAATACTGAAGAGTTCAGCAGTGAG					

Section 81

	(3841)	3841	3850	3860	3870	3888
ClareAJ251507	(3534)	TCAGAACTAGAAGAAAGCAAAGAGAAATTAAATGCAACCAGCTCATCT				
huNall18 (AK)	(3377)	TCAGAACTAGAAGAAAGCAAAGAGAAATTAAATGCAACCAGCTCATCT				
JeongAF225987	(3841)	TCAGAACTAGAAGAAAGCAAAGAGAAATTAAATGCAACCAGCTCATCT				
Consensus	(3841)	TCAGAACTAGAAGAAAGCAAAGAGAAATTAAATGCAACCAGCTCATCT				

Section 82

	(3889)	3889	3900	3910	3920	3936
ClareAJ251507	(3582)	GAAGGAAGCACAGTTGATGTTGTTCTACCCCGAGAAGGTGAACAAGCT				
huNall18 (AK)	(3425)	GAAGGAAGCACAGTTGATGTTGTTCTACCCCGAGAAGGTGAACAAGCT				
JeongAF225987	(3889)	GAAGGAAGCACAGTTGATGTTGTTCTACCCCGAGAAGGTGAACAAGCT				
Consensus	(3889)	GAAGGAAGCACAGTTGATGTTGTTCTACCCCGAGAAGGTGAACAAGCT				

Section 83

	(3937)	3937	3950	3960	3970	3984
ClareAJ251507	(3630)	GAAACTGAACCCGAAGAAGACCTTAAACCGGAAGCTTGTTTTACTGAA				
huNall18 (AK)	(3473)	GAAACTGAACCCGAAGAAGACCTTAAACCGGAAGCTTGTTTTACTGAA				
JeongAF225987	(3937)	GAAACTGAACCCGAAGAAGACTTTAAACCGGAAGCTTGTTTTACTGAA				
Consensus	(3937)	GAAACTGAACCCGAAGAAGACCTTAAACCGGAAGCTTGTTTTACTGAA				

Section 84

	(3985)	3985	3990	4000	4010	4020	4032
ClareAJ251507	(3678)	GGATGTATTAAAAAGTTTCCATTCTGTCAAGTAAGTACAGAAGAAGGC					
huNall18 (AK)	(3521)	GGATGTATTAAAAAGTTTCCATTCTGTCAAGTAAGTACAGAAGAAGGC					
JeongAF225987	(3985)	GGGTGTATTAAAAAGTTTCCATTCTGTCAAGTAAGTACAGAAGAAGGC					
Consensus	(3985)	GGATGTATTAAAAAGTTTCCATTCTGTCAAGTAAGTACAGAAGAAGGC					

Section 85						
	(4033)	4033	4040	4050	4060	4070 4080
ClareAJ251507 (3726)		AAAGGGAAGATCTGGTGGGAATCTTCGAAAAACCTGCTACAGTATTGTT				
huNall18 (AK) (3569)		AAAGGGAAGATCTGGTGGGAATCTTCGAAAAACCTGCTACAGTATTGTT				
JeongAF225987 (4033)		AAAGGGAAGATCTGGTGGGAATCTTCGAAAAACCTGCTACAGTATTGTT				
Consensus (4033)		AAAGGGAAGATCTGGTGGGAATCTTCGAAAAACCTGCTACAGTATTGTT				
Section 86						
	(4081)	4081	4090	4100	4110	4128
ClareAJ251507 (3774)		GAGCACAACTGGTTTGAGACTTTCATTGTGTTTCATGATCCTTCTCAGT				
huNall18 (AK) (3617)		GAGCACAACTGGTTTGAGACTTTCATTGTGTTTCATGATCCTTCTCAGT				
JeongAF225987 (4081)		GAGCACAACTGGTTTGAGACTTTCATTGTGTTTCATGATCCTTCTCAGT				
Consensus (4081)		GAGCACAACTGGTTTGAGACTTTCATTGTGTTTCATGATCCTTCTCAGT				
Section 87						
	(4129)	4129	4140	4150	4160	4176
ClareAJ251507 (3822)		AGTGGTGCATTGGCCTTTGAAGATATATACATTGAACAGCGAAAGACT				
huNall18 (AK) (3665)		AGTGGTGCATTGGCCTTTGAAGATATATACATTGAACAGCGAAAGACT				
JeongAF225987 (4129)		AGTGGTGCATTGGCCTTTGAAGATATATACATTGAACAGCGAAAGACT				
Consensus (4129)		AGTGGTGCATTGGCCTTTGAAGATATATACATTGAACAGCGAAAGACT				
Section 88						
	(4177)	4177	4190	4200	4210	4224
ClareAJ251507 (3870)		ATCAAAACCATGCTAGAAATATGCTGACAAAGTCTTTACCTATATATTC				
huNall18 (AK) (3713)		ATCAAAACCATGCTAGAAATATGCTGACAAAGTCTTTACCTATATATTC				
JeongAF225987 (4177)		ATCAAAACCATGCTAGAAATATGCTGACAAAGTCTTTACCTATATATTC				
Consensus (4177)		ATCAAAACCATGCTAGAAATATGCTGACAAAGTCTTTACCTATATATTC				
Section 89						
	(4225)	4225	4230	4240	4250	4260 4272
ClareAJ251507 (3918)		ATTCTGGAAATGCTTCTCAAATGGGTTGCTTATGGATTTCAAACATAT				
huNall18 (AK) (3761)		ATTCTGGAAATGCTTCTCAAATGGGTTGCTTATGGATTTCAAACATAT				
JeongAF225987 (4225)		ATTCTGGAAATGCTTCTCAAATGGGTTGCTTATGGATTTCAAACATAT				
Consensus (4225)		ATTCTGGAAATGCTTCTCAAATGGGTTGCTTATGGATTTCAAACATAT				
Section 90						
	(4273)	4273	4280	4290	4300	4310 4320
ClareAJ251507 (3966)		TTCACCTAATGCCTGGTGCTGGCTAGATTTCTTGATCGTTGATGTTTCT				
huNall18 (AK) (3809)		TTCACCTAATGCCTGGTGCTGGCTAGATTTCTTGATCGTTGATGTTTCT				
JeongAF225987 (4273)		TTCACCTAATGCCTGGTGCTGGCTAGATTTCTTGATCGTTGATGTTTCT				
Consensus (4273)		TTCACCTAATGCCTGGTGCTGGCTAGATTTCTTGATCGTTGATGTTTCT				
Section 91						
	(4321)	4321	4330	4340	4350	4368
ClareAJ251507 (4014)		TTGGTTAGCCTGGTAGCCAATGCTCTTGGCTACTCAGAACTCGGTGCC				
huNall18 (AK) (3857)		TTGGTTAGCCTGGTAGCCAATGCTCTTGGCTACTCAGAACTCGGTGCC				
JeongAF225987 (4321)		TTGGTTAGCCTGGTAGCCAATGCTCTTGGCTACTCAGAACTCGGTGCC				
Consensus (4321)		TTGGTTAGCCTGGTAGCCAATGCTCTTGGCTACTCAGAACTCGGTGCC				

Section 92

	(4369)	4369	4380	4390	4400	4416
ClareAJ251507	(4062)	ATCAAATCATTACGGACATTAAGAGCTTTAAGACCTCTAAGAGCCTTA				
huNall18 (AK)	(3905)	ATCAAATCATTACGGACATTAAGAGCTTTAAGACCTCTAAGAGCCTTA				
JeongAF225987	(4369)	ATCAAATCATTACGGACATTAAGAGCTTTAAGACCTCTAAGAGCCTTA				
Consensus	(4369)	ATCAAATCATTACGGACATTAAGAGCTTTAAGACCTCTAAGAGCCTTA				

Section 93

	(4417)	4417	4430	4440	4450	4464
ClareAJ251507	(4110)	TCCCGGTTTGAAGGCATGAGGGTGGTTGTGAATGCTCTTGTGGAGCA				
huNall18 (AK)	(3953)	TCCCGGTTTGAAGGCATGAGGGTGGTTGTGAATGCTCTTGTGGAGCA				
JeongAF225987	(4417)	TCCCGGTTTGAAGGCATGAGGGTGGTTGTGAATGCTCTTGTGGAGCA				
Consensus	(4417)	TCCCGGTTTGAAGGCATGAGGGTGGTTGTGAATGCTCTTGTGGAGCA				

Section 94

	(4465)	4465	4470	4480	4490	4500	4512
ClareAJ251507	(4158)	ATTCCCTCTATCATGAATGTGCTGTTGGTCTGTCTCATCTTCTGGTTG					
huNall18 (AK)	(4001)	ATTCCCTCTATCATGAATGTGCTGTTGGTCTGTCTCATCTTCTGGTTG					
JeongAF225987	(4465)	ATTCCCTCTATCATGAATGTGCTGTTGGTCTGTCTCATCTTCTGGTTG					
Consensus	(4465)	ATTCCCTCTATCATGAATGTGCTGTTGGTCTGTCTCATCTTCTGGTTG					

Section 95

	(4513)	4513	4520	4530	4540	4550	4560
ClareAJ251507	(4206)	ATCTTTAGCATCATGGGTGTGAATTTGTTTGCTGGCAAGTTCTACCAC					
huNall18 (AK)	(4049)	ATCTTTAGCATCATGGGTGTGAATTTGTTTGCTGGCAAGTTCTACCAC					
JeongAF225987	(4513)	ATCTTTAGCATCATGGGTGTGAATTTGTTTGCTGGCAAGTTCTACCAC					
Consensus	(4513)	ATCTTTAGCATCATGGGTGTGAATTTGTTTGCTGGCAAGTTCTACCAC					

Section 96

	(4561)	4561	4570	4580	4590	4608
ClareAJ251507	(4254)	TGTGTTAACATGACAACGGGTAAACATGTTTGACATTAGTGATGTTAAC				
huNall18 (AK)	(4097)	TGTGTTAACATGACAACGGGTAAACATGTTTGACATTAGTGATGTTAAC				
JeongAF225987	(4561)	TGTGTTAACATGACAACGGGTAAACATGTTTGACATTAGTGATGTTAAC				
Consensus	(4561)	TGTGTTAACATGACAACGGGTAAACATGTTTGACATTAGTGATGTTAAC				

Section 97

	(4609)	4609	4620	4630	4640	4656
ClareAJ251507	(4302)	AATTTGAGTGACTGTCAGGCTCTTGGCAAGCAAGCTCGGTGGAAAAAC				
huNall18 (AK)	(4145)	AATTTGAGTGACTGTCAGGCTCTTGGCAAGCAAGCTCGGTGGAAAAAC				
JeongAF225987	(4609)	AATTTGAGTGACTGTCAGGCTCTTGGCAAGCAAGCTCGGTGGAAAAAC				
Consensus	(4609)	AATTTGAGTGACTGTCAGGCTCTTGGCAAGCAAGCTCGGTGGAAAAAC				

Section 98

	(4657)	4657	4670	4680	4690	4704
ClareAJ251507	(4350)	GTGAAAGTAACTTTGATAATGTTGGCGCTGGCTATCTTGCACTGCTT				
huNall18 (AK)	(4193)	GTGAAAGTAACTTTGATAATGTTGGCGCTGGCTATCTTGCACTGCTT				
JeongAF225987	(4657)	GTGAAAGTAACTTTGATAATGTTGGCGCTGGCTATCTTGCACTGCTT				
Consensus	(4657)	GTGAAAGTAACTTTGATAATGTTGGCGCTGGCTATCTTGCACTGCTT				

Section 99

	(4705)	4705	4710	4720	4730	4740	4752
ClareAJ251507 (4398)		CAAGTGGCCACATTTAAAGGCTGGATGGATATTATGTATGCAGCTGTT					
huNall18 (AK) (4241)		CAAGTGGCCACATTTAAAGGCTGGATGGATATTATGTATGCAGCTGTT					
JeongAF225987 (4705)		CAAGTGGCCACATTTAAAGGCTGGATGGATATTATGTATGCAGCTGTT					
Consensus (4705)		CAAGTGGCCACATTTAAAGGCTGGATGGATATTATGTATGCAGCTGTT					

Section 100

	(4753)	4753	4760	4770	4780	4790	4800
ClareAJ251507 (4446)		GATTCACGAGATGTTAAACTTCAGCCTGTATATGAAGAAAATCTGTAC					
huNall18 (AK) (4289)		GATTCACGAGATGTTAAACTTCAGCCTGTATATGAAGAAAATCTGTAC					
JeongAF225987 (4753)		GATTCACGAGATGTTAAACTTCAGCCTGTATATGAAGAAAATCTGTAC					
Consensus (4753)		GATTCACGAGATGTTAAACTTCAGCCTGTATATGAAGAAAATCTGTAC					

Section 101

	(4801)	4801	4810	4820	4830	4848
ClareAJ251507 (4494)		ATGTATTTTATACTTTGTCATCTTTATCATCTTTGGGTCATTCTTCACT				
huNall18 (AK) (4337)		ATGTATTTTATACTTTGTCATCTTTATCATCTTTGGGTCATTCTTCACT				
JeongAF225987 (4801)		ATGTATTTTATACTTTGTCATCTTTATCATCTTTGGGTCATTCTTCACT				
Consensus (4801)		ATGTATTTTATACTTTGTCATCTTTATCATCTTTGGGTCATTCTTCACT				

Section 102

	(4849)	4849	4860	4870	4880	4896
ClareAJ251507 (4542)		CTGAATCTATTTCATTGGTGTTCATCATAGATAAACTTCAACCAGCAGAAA				
huNall18 (AK) (4385)		CTGAATCTATTTCATTGGTGTTCATCATAGATAAACTTCAACCAGCAGAAA				
JeongAF225987 (4849)		CTGAATCTATTTCATTGGTGTTCATCATAGATAAACTTCAACCAGCAGAAA				
Consensus (4849)		CTGAATCTATTTCATTGGTGTTCATCATAGATAAACTTCAACCAGCAGAAA				

Section 103

	(4897)	4897	4910	4920	4930	4944
ClareAJ251507 (4590)		AAGAAGTTTGGAGGTCAAGACATCTTTATGACAGAGGAACAGAAAAAA				
huNall18 (AK) (4433)		AAGAAGTTTGGAGGTCAAGACATCTTTATGACAGAGGAACAGAAAAAA				
JeongAF225987 (4897)		AAGAAGTTTGGAGGTCAAGACATCTTTATGACAGAGGAACAGAAAAAA				
Consensus (4897)		AAGAAGTTTGGAGGTCAAGACATCTTTATGACAGAGGAACAGAAAAAA				

Section 104

	(4945)	4945	4950	4960	4970	4980	4992
ClareAJ251507 (4638)		TATTACAATGCAATGAAGAACTTGGATCCAAGAAACCTCAGAAACCC					
huNall18 (AK) (4481)		TATTACAATGCAATGAAGAACTTGGATCCAAGAAACCTCAGAAACCC					
JeongAF225987 (4945)		TATTACAATGCAATGAAGAACTTGGATCCAAGAAACCTCAGAAACCC					
Consensus (4945)		TATTACAATGCAATGAAGAACTTGGATCCAAGAAACCTCAGAAACCC					

Section 105

	(4993)	4993	5000	5010	5020	5030	5040
ClareAJ251507 (4686)		ATACCTCGCCCAGCAAACAAATTCCAAGGAATGGTCTTTGATTTTGTA					
huNall18 (AK) (4529)		ATACCTCGCCCAGCAAACAAATTCCAAGGAATGGTCTTTGATTTTGTA					
JeongAF225987 (4993)		ATACCTCGCCCAGCAAACAAATTCCAAGGAATGGTCTTTGATTTTGTA					
Consensus (4993)		ATACCTCGCCCAGCAAACAAATTCCAAGGAATGGTCTTTGATTTTGTA					

Section 106							
	(5041)	5041	5050	5060	5070	5088	
ClareAJ251507	(4734)	ACCAGACAAGTCTTTGATATCAGCATCATGATCCTCATCTGCCTCAAC					
huNall18 (AK)	(4577)	ACCAGACAAGTCTTTGATATCAGCATCATGATCCTCATCTGCCTCAAC					
JeongAF225987	(5041)	ACCAGACAAGTCTTTGATATCAGCATCATGATCCTCATCTGCCTCAAC					
Consensus	(5041)	ACCAGACAAGTCTTTGATATCAGCATCATGATCCTCATCTGCCTCAAC					
Section 107							
	(5089)	5089	5100	5110	5120	5136	
ClareAJ251507	(4782)	ATGGTCACCATGATGGTGGAAACGGATGACCAGGGCAAATACATGACC					
huNall18 (AK)	(4625)	ATGGTCACCATGATGGTGGAAACGGATGACCAGGGCAAATACATGACC					
JeongAF225987	(5089)	ATGGTCACCATGATGGTGGAAACGGATGACCAGGGCAAATACATGACC					
Consensus	(5089)	ATGGTCACCATGATGGTGGAAACGGATGACCAGGGCAAATACATGACC					
Section 108							
	(5137)	5137	5150	5160	5170	5184	
ClareAJ251507	(4830)	CTAGTTTTGTCCCGGATCAACCTAGTGTTTCATTGTTCTGTTCACTGGA					
huNall18 (AK)	(4673)	CTAGTTTTGTCCCGGATCAACCTAGTGTTTCATTGTTCTGTTCACTGGA					
JeongAF225987	(5137)	CTAGTTTTGTCCCGGATCAACCTAGTGTTTCATTGTTCTGTTCACTGGA					
Consensus	(5137)	CTAGTTTTGTCCCGGATCAACCTAGTGTTTCATTGTTCTGTTCACTGGA					
Section 109							
	(5185)	5185	5190	5200	5210	5220	5232
ClareAJ251507	(4878)	GAATTTGTGCTGAAGCTCGTTTCCCTCAGACACTACTACTTCACTATA					
huNall18 (AK)	(4721)	GAATTTGTGCTGAGGCTCGTTTCCCTCAGACACTACTACTTCACTATA					
JeongAF225987	(5185)	GAATTTGTGCTGAAGCTCGTTTCCCTCAGACACTACTACTTCACTATA					
Consensus	(5185)	GAATTTGTGCTGAAGCTCGTCTCCCTCAGACACTACTACTTCACTATA					
Section 110							
	(5233)	5233	5240	5250	5260	5270	5280
ClareAJ251507	(4926)	GGCTGGAACATCTTTGACTTTGTGGTGGTGATTCTCTCCATTGTAGGT					
huNall18 (AK)	(4769)	GGCTGGAACATCTTTGACTTTGTGGTGGTGATTCTCTCCATTGTAGGT					
JeongAF225987	(5233)	GGCTGGAACATCTTTGACTTTGTGGTGGTGATTCTCTCCATTGTAGGT					
Consensus	(5233)	GGCTGGAACATCTTTGACTTTGTGGTGGTGATTCTCTCCATTGTAGGT					
Section 111							
	(5281)	5281	5290	5300	5310	5328	
ClareAJ251507	(4974)	ATGTTTCTGGCTGAGATGATAGAAAAGTATTCTGTGTCCCCTACCTTG					
huNall18 (AK)	(4817)	ATGTTTCTGGCTGAGATGATAGAAAAGTATTCTGTGTCCCCTACCTTG					
JeongAF225987	(5281)	ATGTTTCTGGCTGAGATGATAGAAAAGTATTCTGTGTCCCCTACCTTG					
Consensus	(5281)	ATGTTTCTGGCTGAGATGATAGAAAAGTATTTTGTGTCCCCTACCTTG					
Section 112							
	(5329)	5329	5340	5350	5360	5376	
ClareAJ251507	(5022)	TTCCGAGTGATCCGTCTTGCCAGGATTGGCCGAATCCTACGTCTGATC					
huNall18 (AK)	(4865)	TTCCGAGTGATCCGTCTTGCCAGGATTGGCCGAATCCTACGTCTGATC					
JeongAF225987	(5329)	TTCCGAGTGATCCGTCTTGCCAGGATTGGCCGAATCCTACGTCTGATC					
Consensus	(5329)	TTCCGAGTGATCCGTCTTGCCAGGATTGGCCGAATCCTACGTCTGATC					

Section 113

	(5377)	5377	5390	5400	5410	5424
ClareAJ251507 (5070)		AAAGGAGCAAAGGGGATCCGCACGCTGCTCTTTGCTTTGATGATGTCC				
huNall18 (AK) (4913)		AAAGGAGCAAAGGGGATCCGCACGCTGCTCTTTGCTTTGATGATGTCC				
JeongAF225987 (5377)		AAAGGAGCAAAGGGGATCCGCACGCTGCTCTTTGCTTTGATGATGTCC				
Consensus (5377)		AAAGGAGCAAAGGGGATCCGCACGCTGCTCTTTGCTTTGATGATGTCC				

Section 114

	(5425)	5425	5430	5440	5450	5460	5472
ClareAJ251507 (5118)		CTTCCTGCGTTGTTTAAACATCGGCCTCCTGCTCTTCCTGGTCATGTTT					
huNall18 (AK) (4961)		CTTCCTGCGTTGTTTAAACATCGGCCTCCTGCTCTTCCTGGTCATGTTT					
JeongAF225987 (5425)		CTTCCTGCGTTGTTTAAACATCGGCCTCCTGCTCTTCCTGGTCATGTTT					
Consensus (5425)		CTTCCTGCGTTGTTTAAACATCGGCCTCCTGCTCTTCCTGGTCATGTTT					

Section 115

	(5473)	5473	5480	5490	5500	5510	5520
ClareAJ251507 (5166)		ATCTATGCCATCTTTGGGATGTCCAACCTTTGCCTATGTTAAAAAGGAA					
huNall18 (AK) (5009)		ATCTATGCCATCTTTGGGATGTCCAACCTTTGCCTATGTTAAAAAGGAA					
JeongAF225987 (5473)		ATCTATGCCATCTTTGGGATGTCCAACCTTTGCCTATGTTAAAAAGGAA					
Consensus (5473)		ATCTATGCCATCTTTGGGATGTCCAACCTTTGCCTATGTTAAAAAGGAA					

Section 116

	(5521)	5521	5530	5540	5550	5568
ClareAJ251507 (5214)		GCTGGAATTGATGACATGTTCAACTTTGAGACCTTTGGCAACAGCATG				
huNall18 (AK) (5057)		GCTGGAATTGATGACATGTTCAACTTTGAGACCTTTGGCAACAGCATG				
JeongAF225987 (5521)		GCTGGAATTGATGACATGTTCAACTTTGAGACCTTTGGCAACAGCATG				
Consensus (5521)		GCTGGAATTGATGACATGTTCAACTTTGAGACCTTTGGCAACAGCATG				

Section 117

	(5569)	5569	5580	5590	5600	5616
ClareAJ251507 (5262)		ATCTGCTTGTTCCAAATTACAACCTCTGCTGGCTGGGATGGATTGCTA				
huNall18 (AK) (5105)		ATCTGCTTGTTCCAAATTACAACCTCTGCTGGCTGGGATGGATTGCTA				
JeongAF225987 (5569)		ATCTGCTTGTTCCAAATTACAACCTCTGCTGGCTGGGATGGATTGCTA				
Consensus (5569)		ATCTGCTTGTTCCAAATTACAACCTCTGCTGGCTGGGATGGATTGCTA				

Section 118

	(5617)	5617	5630	5640	5650	5664
ClareAJ251507 (5310)		GCACCTATTCTTAATAGTGCACCACCCGACTGTGACCCTGACACAATT				
huNall18 (AK) (5153)		GCACCTATTCTTAATAGTGCACCACCCGACTGTGACCCTGACACAATT				
JeongAF225987 (5617)		GCACCTATTCTTAATAGTGCACCACCCGACTGTGACCCTGACACAATT				
Consensus (5617)		GCACCTATTCTTAATAGTGCACCACCCGACTGTGACCCTGACACAATT				

Section 119

	(5665)	5665	5670	5680	5690	5700	5712
ClareAJ251507 (5358)		CACCCTGGCAGCTCAGTTAAGGGAGACCGTGGGACCCATCTGTTGGG					
huNall18 (AK) (5201)		CACCCTGGCAGCTCAGTTAAGGGAGACCGTGGGACCCATCTGTTGGG					
JeongAF225987 (5665)		CACCCTGGCAGCTCAGTTAAGGGAGACCGTGGGACCCATCTGTTGGG					
Consensus (5665)		CACCCTGGCAGCTCAGTTAAGGGAGACTGTGGGAACCCATCTGTTGGG					

Section 120						
	(5713)	5713	5720	5730	5740	5750 5760
ClareAJ251507 (5406)		ATTTTCTTTTTTCGTCAGTTACATCATCATATCCTTCCTGGTTGTGGTG				
huNall18 (AK) (5249)		ATTTTCTTTTTTCGTCAGTTACATCATCATATCCTTCCTGGTTGTGGTG				
JeongAF225987 (5713)		ATTTTCTTTTTTCGTCAGTTACATCATCATATCCTTCCTGGTTGTGGTG				
Consensus (5713)		ATTTTCTTTTTTCGTCAGTTACATCATCATATCCTTCCTGGTTGTGGTG				
Section 121						
	(5761)	5761	5770	5780	5790	5808
ClareAJ251507 (5454)		AACATGTACATCGCGGTCATCCTGGAGAACTTCAGTGTGCTACTGAA				
huNall18 (AK) (5297)		AACATGTACATCGCGGTCATCCTGGAGAACTTCAGTGTGCTACTGAA				
JeongAF225987 (5761)		AACATGTACATCGCGGTCATCCTGGAGAACTTCAGTGTGCTACTGAA				
Consensus (5761)		AACATGTACATCGCGGTCATCCTGGAGAACTTCAGTGTGCTACTGAA				
Section 122						
	(5809)	5809	5820	5830	5840	5856
ClareAJ251507 (5502)		GAAAGTGCAGAGCCCCTGAGTGAGGATGACTTTGAGATGTTCTATGAG				
huNall18 (AK) (5345)		GAAAGTGCAGAGCCCCTGAGTGAGGATGACTTTGAGATGTTCTATGAG				
JeongAF225987 (5809)		GAAAGTGCAGAGCCCCTGAGTGAGGATGACTTTGAGATGTTCTATGAG				
Consensus (5809)		GAAAGTGCAGAGCCCCTGAGTGAGGATGACTTTGAGATGTTCTATGAG				
Section 123						
	(5857)	5857	5870	5880	5890	5904
ClareAJ251507 (5550)		GTTTGGGAAAAGTTTGATCCCGATGCGACCCAGTTTATAGAGTTCTCT				
huNall18 (AK) (5393)		GTTTGGGAAAAGTTTGATCCCGATGCGACCCAGTTTATAGAGTTCTCT				
JeongAF225987 (5857)		GTTTGGGAAAAGTTTGATCCCGATGCGACCCAGTTTATAGAGTTCTCT				
Consensus (5857)		GTTTGGGAAAAGTTTGATCCCGATGCGACCCAGTTTATAGAGTTCTCT				
Section 124						
	(5905)	5905	5910	5920	5930	5940 5952
ClareAJ251507 (5598)		AAACTCTCTGATTTTGCAGCTGCCCTGGATCCTCCTCTTCTCATAGCA				
huNall18 (AK) (5441)		AAACTCTCTGATTTTGCAGCTGCCCTGGATCCTCCTCTTCTCATAGCA				
JeongAF225987 (5905)		AAACTCTCTGATTTTGCAGCTGCCCTGGATCCTCCTCTTCTCATAGCA				
Consensus (5905)		AAACTCTCTGATTTTGCAGCTGCCCTGGATCCTCCTCTTCTCATAGCA				
Section 125						
	(5953)	5953	5960	5970	5980	5990 6000
ClareAJ251507 (5646)		AAACCCAACAAAGTCCAGCTTATTGCCATGGATCTGCCCATGGTCAGT				
huNall18 (AK) (5489)		AAACCCAACAAAGTCCAGCTTATTGCCATGGATCTGCCCATGGTCAGT				
JeongAF225987 (5953)		AAACCCAACAAAGTCCAGCTTATTGCCATGGATCTGCCCATGGTCAGT				
Consensus (5953)		AAACCCAACAAAGTCCAGCTTATTGCCATGGATCTGCCCATGGTCAGT				
Section 126						
	(6001)	6001	6010	6020	6030	6048
ClareAJ251507 (5694)		GGTGACCGGATCCACTGTCTTGATATTTTATTTGCCTTTACAAAGCGT				
huNall18 (AK) (5537)		GGTGACCGGATCCACTGTCTTGATATTTTATTTGCCTTTACAAAGCGT				
JeongAF225987 (6001)		GGTGACCGGATCCACTGTCTTGATATTTTATTTGCCTTTACAAAGCGT				
Consensus (6001)		GGTGACCGGATCCACTGTCTTGATATTTTATTTGCCTTTACAAAGCGT				

Section 127

	(6049)	6049	6060	6070	6080	6096
ClareAJ251507 (5742)		GTTTTG	GGTGAGAGTGGAGAGATGGATGCCCTTCGAATACAGATGGAA			
huNall18 (AK) (5585)		GTTTTG	GGTGAGAGTGGAGAGATGGATGCCCTTCGAATACAGATGGAA			
JeongAF225987 (6049)		GTTTTG	GGTGAGAGTGGAGAGATGGATGCCCTTCGAATACAGATGGAA			
Consensus (6049)		GTTTTG	GGTGAGAGTGGAGAGATGGATGCCCTTCGAATACAGATGGAA			

Section 128

	(6097)	6097	6110	6120	6130	6144
ClareAJ251507 (5790)		GACAGGTTTATGGCATCAAACCCCTCCAAAGTCTCTTATGAGCCTATT				
huNall18 (AK) (5633)		GACAGGTTTATGGCATCAAACCCCTCCAAAGTCTCTTATGAGCCTATT				
JeongAF225987 (6097)		GACAGGTTTATGGCATCAAACCCCTCCAAAGTCTCTTATGAGCCTATT				
Consensus (6097)		GACAGGTTTATGGCATCAAACCCCTCCAAAGTCTCTTATGAGCCTATT				

Section 129

	(6145)	6145	6150	6160	6170	6180	6192
ClareAJ251507 (5838)		ACAACCACCTTTGAAACGTAAACAAGAGGAGGTGTCTGCCGCTATCATT					
huNall18 (AK) (5681)		ACAACCACCTTTGAAACGTAAACAAGAGGAGGTGTCTGCCGCTATCATT					
JeongAF225987 (6145)		ACAACCACCTTTGAAACGTAAACAAGAGGAGGTGTCTGCCGCTATCATT					
Consensus (6145)		ACAACCACCTTTGAAACGTAAACAAGAGGAGGTGTCTGCCGCTATCATT					

Section 130

	(6193)	6193	6200	6210	6220	6230	6240
ClareAJ251507 (5886)		CAGCGTAATTTTCAGATGTTATCTTTTAAAGCAAAGGTTAAAAAATATA					
huNall18 (AK) (5729)		CAGCGTAATTTTCAGATGTTATCTTTTAAAGCAAAGGTTAAAAAATATA					
JeongAF225987 (6193)		CAGCGTAATTTTCAGATGTTATCTTTTAAAGCAAAGGTTAAAAAATATA					
Consensus (6193)		CAGCGTAATTTTCAGATGTTATCTTTTAAAGCAAAGGTTAAAAAATATA					

Section 131

	(6241)	6241	6250	6260	6270	6288
ClareAJ251507 (5934)		TCAAGTAACTATAACAAAGAGGCAATTAAAGGGAGGATTGACTTACCT				
huNall18 (AK) (5777)		TCAAGTAACTATAACAAAGAGGCAATTAAAGGGAGGATTGACTTACCT				
JeongAF225987 (6241)		TCAAGTAACTATAACAAAGAGGCAATTAAAGGGAGGATTGACTTACCT				
Consensus (6241)		TCAAGTAACTATAACAAAGAGGCAATTAAAGGGAGGATTGACTTACCT				

Section 132

	(6289)	6289	6300	6310	6320	6336
ClareAJ251507 (5982)		ATAAAACAAGACATGATTATTGACAAACTAAATGGGAACCTCCACTCCA				
huNall18 (AK) (5825)		ATAAAACAAGACATGATTATTGACAAACTAAATGGGAACCTCCACTCCA				
JeongAF225987 (6289)		ATAAAACAAGACATGATTATTGACAAACTAAATGGGAACCTCCACTCCA				
Consensus (6289)		ATAAAACAAGACATGATTATTGACAAACTAAATGGGAACCTCCACTCCA				

Section 133

	(6337)	6337	6350	6360	6370	6384
ClareAJ251507 (6030)		GAAAAAACAGATGGGAGTTCCTCTACCACCTCTCCTCCTTCCTATGAT				
huNall18 (AK) (5873)		GAAAAAACAGATGGGAGTTCCTCTACCACCTCTCCTCCTTCCTATGAT				
JeongAF225987 (6337)		GAAAAAACAGATGGGAGTTCCTCTACCACCTCTCCTCCTTCCTATGAT				
Consensus (6337)		GAAAAAACAGATGGGAGTTCCTCTACCACCTCTCCTCCTTCCTATGAT				

Section 134						
	(6385)	6385	6390	6400	6410	6420 6432
ClareAJ251507 (6078)		AGTGTAAACAAAACCAGACAAGGAAAAGTTTGAGAAAGACAAACCAGAA				
huNall18 (AK) (5921)		AGTGTAAACAAAACCAGACAAGGAAAAGTTTGAGAAAGACAAACCAGAA				
JeongAF225987 (6385)		AGTGTAAACAAAACCAGACAAGGAAAAGTTTGAGAAAGACAAACCAGAA				
Consensus (6385)		AGTGTAAACAAAACCAGACAAGGAAAAGTTTGAGAAAGACAAACCAGAA				
Section 135						
	(6433)	6433	6440	6450	6460	6470 6480
ClareAJ251507 (6126)		AAAGAAAGCAAAGGAAAAGAGGTCAGAGAAAATCAAAAGTAAAAAGAA				
huNall18 (AK) (5969)		AAAGAAAGCAAAGGAAAAGAGGTCAGAGAAAATCAAAAGTAAAAAGAA				
JeongAF225987 (6433)		AAAGAAAGCAAAGGAAAAGAGGTCAGAGAAAATCAAAAGTAAAAAGAA				
Consensus (6433)		AAAGAAAGCAAAGGAAAAGAGGTCAGAGAAAATCAAAAGTAAAAAGAA				
Section 136						
	(6481)	6481	6490	6500	6510	6528
ClareAJ251507 (6174)		ACAAAGAATTATCTTTGTGATCAATTGTTTACAGCCTATGAAGGTAAA				
huNall18 (AK) (6017)		ACAAAGAATTATCTTTGTGATCAATTGTTTACAGCCTATGAAGGTAAA				
JeongAF225987 (6481)		ACAAAGAATTATCTTTGTGATCAATTGTTTACAGCCTATGAAGGTAAA				
Consensus (6481)		ACAAAGAATTATCTTTGTGATCAATTGTTTACAGCCTATGAAGGTAAA				
Section 137						
	(6529)	6529	6540	6550	6560	6576
ClareAJ251507 (6222)		GTATATGTGTCAACTGGACTTCAAG AGGAGGTCCATGCCAAACTGACT				
huNall18 (AK) (6065)		GTATATGTGTCAACTGGACTTCAAG -----				
JeongAF225987 (6529)		GTATATGTGTCAACTGGACTTCAAG AGGAGGTCCATGCCAAACTGACT				
Consensus (6529)		GTATATGTGTCAACTGGACTTCAAGAGGAGGTCCATGCCAAACTGACT				
Section 138						
	(6577)	6577	6590	6600	6610	6624
ClareAJ251507 (6270)		GTATTAACTAATACCTATAGTGAGTGCCTATACAGACAGTGAAGTGA				
huNall18 (AK) (6090)		-----				
JeongAF225987 (6577)		GTATTAACTAATACCTATAGTGAGTGCCTATACAGACAGTGAAGTGA				
Consensus (6577)		GTTTTAACAATACTCATAGTCAGTGCCTATACAGACAGTGAAGTGA				
Section 139						
	(6625)	6625	6630	6640	6650	6660 6672
ClareAJ251507 (6318)		CCTCTCTGTCACTGCAACTCTGTGAAGCAGGGTATCAAC TTGACAAG				
huNall18 (AK) (6090)		-----				
JeongAF225987 (6625)		CCTCTCTGTCACTGCAACTCTGTGAAGCAGGGTATCAAC TTGACAAG				
Consensus (6625)		CCTCTCTGTCACTGCAACTCTGTGAAGCAGGGTATCAAC TTGACAAG				
Section 140						
	(6673)	6673	6680	6690	6700	6710 6720
ClareAJ251507 (6366)		AGGTTGCTGTTTTTATTACCAGCTGACACTGCTGAGGAGAAACCCAAT				
huNall18 (AK) (6090)		-----				
JeongAF225987 (6673)		AGGTTGCTGTTTTTATTACCAGCTGACACTGCTGAGGAGAAACCCAAT				
Consensus (6673)		AGGTTGCTGTTTTTATTACCAGCTGACACTGCTGAGGAGAAACCCAAT				

Section 141						
	(6721)	6721	6730	6740	6750	6768
ClareAJ251507	(6414)	GGCTACCTAGACTATAGGGATAGTTGCTGC AAAAGTGAACATTGTAAC TA				
huNall18 (AK)	(6090)	-----				
JeongAF225987	(6721)	GGCTACCTAGACTATAGGGATAGTTGCTGC AAAAGTGAACATTGTAAC TA				
Consensus	(6721)	GGCTACCTAGACTATAGGGATAGTTGCTGC AAAAGTGAACATTGTAAC TA				
Section 142						
	(6769)	6769	6780	6790	6800	6816
ClareAJ251507	(6462)	CACCAAACACCTTTAGTACAGTCCTTGCATCCATTCTATTTT TAACTT				
huNall18 (AK)	(6090)	-----				
JeongAF225987	(6769)	CACCAAACACCTTTAGTACAGTCCTTGCATCCATTCTATTTT TAACTT				
Consensus	(6769)	CACCAAACACCTTTAGTACAGTCCTTGCATCCATTCTATTTT TAACTT				
Section 143						
	(6817)	6817	6830	6840	6850	6864
ClareAJ251507	(6510)	CCATATCTGCCATATTTT TACAAAATTTGTTCTAGTGCATTTCCATGG				
huNall18 (AK)	(6090)	-----				
JeongAF225987	(6817)	CCATATCTGCCATATTTT TACAAAATTTGTTCTAGTGCATTTCCATGG				
Consensus	(6817)	CCATATCTGCCATATTTT TACAAAATTTGTTCTAGTGCATTTCCATGG				
Section 144						
	(6865)	6865	6870	6880	6890	6912
ClareAJ251507	(6558)	TCCCAATTTCATAGTTTATTTCATAATGCTATGTCACTATTTT				
huNall18 (AK)	(6090)	-----				
JeongAF225987	(6865)	TCCCAATTTCATAGTTTATTTCATAATGCTATGTCACTATTTT				
Consensus	(6865)	TCCCAATTTCATAGTTTATTTCATAATGCTATGTCACTATTTT				
Section 145						
	(6913)	6913	6920	6930	6940	6960
ClareAJ251507	(6600)	-----				
huNall18 (AK)	(6090)	-----				
JeongAF225987	(6913)	TGAGGTTTACGTTGAAGAAACAGTATACAAGAACCCTGTCTCTCAAAT				
Consensus	(6913)	TGAGGTTTACGTTGAAGAAACAGTATACAAGAACCCTGTCTCTCAAAT				
Section 146						
	(6961)	6961	6970	6980	6990	7008
ClareAJ251507	(6600)	-----				
huNall18 (AK)	(6090)	-----				
JeongAF225987	(6961)	GATCAGACAAAGGTGTTTTGCCAGAGAGATAAAATTTT TGCTCAAAAC				
Consensus	(6961)	GATCAGACAAAGGTGTTTTGCCAGAGAGATAAAATTTT TGCTCAAAAC				
Section 147						
	(7009)	7009	7020	7030	7040	7056
ClareAJ251507	(6600)	-----				
huNall18 (AK)	(6090)	-----				
JeongAF225987	(7009)	CAGAAAAAGAATTGTAATGGCTACAGTTTCAGTTACTTCCATTTTCTA				
Consensus	(7009)	CAGAAAAAGAATTGTAATGGCTACAGTTTCAGTTACTTCCATTTTCTA				

Section 148						
(7057)	7057		7070	7080	7090	7104
ClareAJ251507 (6600)	-----		-----	-----	-----	-----
huNall18 (AK) (6090)	-----		-----	-----	-----	-----
JeongAF225987 (7057)	GATGGCTTTAATTTTGAAAGTATTTTAGTCTGTTATGTTTGTTCCTAT					
Consensus (7057)						
Section 149						
(7105)	7105	7110	7120	7130	7140	7152
ClareAJ251507 (6600)	-----	-----	-----	-----	-----	-----
huNall18 (AK) (6090)	-----	-----	-----	-----	-----	-----
JeongAF225987 (7105)	CTGAACAGTTATGTGCCTGTAAAGTCTCCTCTAATATTTAAAGGATTA					
Consensus (7105)						
Section 150						
(7153)	7153	7160	7170	7180	7190	7200
ClareAJ251507 (6600)	-----	-----	-----	-----	-----	-----
huNall18 (AK) (6090)	-----	-----	-----	-----	-----	-----
JeongAF225987 (7153)	TTTTTATGCAAAGTATTCTGTTTCAGCAAGTGCAAATTTTATTCTAAG					
Consensus (7153)						
Section 151						
(7201)	7201	7210	7220	7230		7248
ClareAJ251507 (6600)	-----	-----	-----	-----	-----	-----
huNall18 (AK) (6090)	-----	-----	-----	-----	-----	-----
JeongAF225987 (7201)	TTTCAGAGCTCTATATTTAATTTAGGTCAAATGCTTTCCAAAAGTAA					
Consensus (7201)						
Section 152						
(7249)	7249	7260	7270	7280		7296
ClareAJ251507 (6600)	-----	-----	-----	-----	-----	-----
huNall18 (AK) (6090)	-----	-----	-----	-----	-----	-----
JeongAF225987 (7249)	TCTAATAAATCCATTCTAGAAAAATATATCTAAAGTATTGCTTTAGAA					
Consensus (7249)						
Section 153						
(7297)	7297	7310	7320	7330		7344
ClareAJ251507 (6600)	-----	-----	-----	-----	-----	-----
huNall18 (AK) (6090)	-----	-----	-----	-----	-----	-----
JeongAF225987 (7297)	TAGTTGTTCCACTTTCTGCTGCAGTATTGCTTTGCCATCTTCTGCTCT					
Consensus (7297)						
Section 154						
(7345)	7345	7350	7360	7370	7380	7392
ClareAJ251507 (6600)	-----	-----	-----	-----	-----	-----
huNall18 (AK) (6090)	-----	-----	-----	-----	-----	-----
JeongAF225987 (7345)	CAGCAAAGCTGATAGTCTATGTCAATTAAATACCCTATGTTATGTAAA					
Consensus (7345)						

							Section 155
(7393)	7393	7400	7410	7420	7430	7440	
ClareAJ251507 (6600)	-----						
huNall118 (AK) (6090)	-----						
JeongAF225987 (7393)	TAGTTATTTTATCCTGTGGTGCATGTTTGGGCAAATATATATATAGCC						
Consensus (7393)							
							Section 156
(7441)	7441	7450	7460	7470		7488	
ClareAJ251507 (6600)	-----						
huNall118 (AK) (6090)	-----						
JeongAF225987 (7441)	TGATAAACAACTTCTATTAAATCAAATATGTACCACAGTGTATGTGTC						
Consensus (7441)							
							Section 157
(7489)	7489	7500	7510	7520		7536	
ClareAJ251507 (6600)	-----						
huNall118 (AK) (6090)	-----						
JeongAF225987 (7489)	TTTGTGCAAGCTTCCAACAGGGATGTATCCTGTATCATTCATTAAACAT						
Consensus (7489)							
							Section 158
(7537)	7537	7550	7560	7570		7584	
ClareAJ251507 (6600)	-----						
huNall118 (AK) (6090)	-----						
JeongAF225987 (7537)	AGTTTAAAGGCTATCACTAATGCATGTTAATATTGCCTATGCTGCTCT						
Consensus (7537)							
							Section 159
(7585)	7585	7590	7600	7610	7620	7632	
ClareAJ251507 (6600)	-----						
huNall118 (AK) (6090)	-----						
JeongAF225987 (7585)	ATTTTACTCAATCCATTCTTCACAAGTCTTGGTTAAAGAATGTCACAT						
Consensus (7585)							
							Section 160
(7633)	7633	7640	7650	7660	7670	7680	
ClareAJ251507 (6600)	-----						
huNall118 (AK) (6090)	-----						
JeongAF225987 (7633)	ATTGGTGATAGAATGAATTCAACCTGCTCTGTCCATTATGTCAAGCAG						
Consensus (7633)							
							Section 161
(7681)	7681	7690	7700	7710		7728	
ClareAJ251507 (6600)	-----						
huNall118 (AK) (6090)	-----						
JeongAF225987 (7681)	AATAATTTGAAGCTATTTACAAACACCTTTACTTTTGCACTTTTAATT						
Consensus (7681)							

						Section 162	
	(7729)	7729	7740	7750	7760	7776	
ClareAJ251507	(6600)	-----	-----	-----	-----	-----	
huNall18 (AK)	(6090)	-----	-----	-----	-----	-----	
JeongAF225987	(7729)	CAACATGAGTATCATATGGTATCTCTCTGGATTTC	AAGGAAACACACT				
Consensus	(7729)						
						Section 163	
	(7777)	7777	7790	7800	7810	7824	
ClareAJ251507	(6600)	-----	-----	-----	-----	-----	
huNall18 (AK)	(6090)	-----	-----	-----	-----	-----	
JeongAF225987	(7777)	GGATACTGCCTACTGACAAAACCTATTCTTCATATTTTGCTAAAAATA					
Consensus	(7777)						
						Section 164	
	(7825)	7825	7830	7840	7850	7860	7872
ClareAJ251507	(6600)	-----	-----	-----	-----	-----	-----
huNall18 (AK)	(6090)	-----	-----	-----	-----	-----	-----
JeongAF225987	(7825)	TGTCTAAAACTTGTTTAAATATAAATAATGTAAAAATATAATCAACTT					
Consensus	(7825)						
						Section 165	
	(7873)	7873	7880	7890	7900	7910	7920
ClareAJ251507	(6600)	-----	-----	-----	-----	-----	-----
huNall18 (AK)	(6090)	-----	-----	-----	-----	-----	-----
JeongAF225987	(7873)	TATTTGTCAGCATTTTGTACATAAGAAAATTATTTTCAGGTTGATGAC					
Consensus	(7873)						
						Section 166	
	(7921)	7921	7930	7940	7950		7968
ClareAJ251507	(6600)	-----	-----	-----	-----	-----	-----
huNall18 (AK)	(6090)	-----	-----	-----	-----	-----	-----
JeongAF225987	(7921)	ATCACAATTTATTTTACTTTATGCTTTTGCTTTTGATTTTAAATCACA					
Consensus	(7921)						
						Section 167	
	(7969)	7969	7980	7990	8000		8016
ClareAJ251507	(6600)	-----	-----	-----	-----	-----	-----
huNall18 (AK)	(6090)	-----	-----	-----	-----	-----	-----
JeongAF225987	(7969)	ATTCCAAACTTTTGAATCCATAAGATTTTCAATGGATAATTTCTCTAA					
Consensus	(7969)						
						Section 168	
	(8017)	8017	8030	8040	8050		8064
ClareAJ251507	(6600)	-----	-----	-----	-----	-----	-----
huNall18 (AK)	(6090)	-----	-----	-----	-----	-----	-----
JeongAF225987	(8017)	AATAAAAGTTAGATAATGGGTTTTATGGATTTCTTTGTTATAATATAT					
Consensus	(8017)						

							Section 169
	(8065)	8065	8070	8080	8090	8100	8112
ClareAJ251507 (6600)		-----	-----	-----	-----	-----	-----
huNall18 (AK) (6090)		-----	-----	-----	-----	-----	-----
JeongAF225987 (8065)		TTTCTACCATTCCAATAGGAGATACATTGGTCAAACACTCAAACCTAG					
Consensus (8065)							
							Section 170
	(8113)	8113	8120	8130	8140	8150	8160
ClareAJ251507 (6600)		-----	-----	-----	-----	-----	-----
huNall18 (AK) (6090)		-----	-----	-----	-----	-----	-----
JeongAF225987 (8113)		ATCATTTTCTACCAACTATGGTTGCCTCAATATAACCTTTTATTCATA					
Consensus (8113)							
							Section 171
	(8161)	8161	8170	8180	8190		8208
ClareAJ251507 (6600)		-----	-----	-----	-----	-----	-----
huNall18 (AK) (6090)		-----	-----	-----	-----	-----	-----
JeongAF225987 (8161)		GATGTTTTTTTTTATTCAACTTTTGTAGTATTTACGTATGCAGACTAG					
Consensus (8161)							
							Section 172
	(8209)	8209	8220	8230	8240		8256
ClareAJ251507 (6600)		-----	-----	-----	-----	-----	-----
huNall18 (AK) (6090)		-----	-----	-----	-----	-----	-----
JeongAF225987 (8209)		TCTTATTTTTTTTAATTCTGCTGCACTAAAGCTATTACAAATATAACA					
Consensus (8209)							
							Section 173
	(8257)	8257	8270	8280	8290		8304
ClareAJ251507 (6600)		-----	-----	-----	-----	-----	-----
huNall18 (AK) (6090)		-----	-----	-----	-----	-----	-----
JeongAF225987 (8257)		TGGACTTTGTTCTTTTGTAGCCATGAACAAAGTGGCAAAGTTGTGCAAT					
Consensus (8257)							
							Section 174
	(8305)	8305	8310	8320	8330	8340	8352
ClareAJ251507 (6600)		-----	-----	-----	-----	-----	-----
huNall18 (AK) (6090)		-----	-----	-----	-----	-----	-----
JeongAF225987 (8305)		TACCTAACATGATATAAATTTTGTGTTTTTGCACAAACCAAAGTTTA					
Consensus (8305)							
							Section 175
	(8353)	8353	8360	8370	8380	8390	8400
ClareAJ251507 (6600)		-----	-----	-----	-----	-----	-----
huNall18 (AK) (6090)		-----	-----	-----	-----	-----	-----
JeongAF225987 (8353)		ATGTTAATTCTTTTACAAAACCTATTTACTGTAGTGTATTGAAGAAGT					
Consensus (8353)							

Section 176					
	(8401)	8401	8410	8420	8430
ClareAJ251507 (6600)		-----	-----	-----	-----
huNall18 (AK) (6090)		-----	-----	-----	-----
JeongAF225987 (8401)		GCATGCAGGGAATTGCTATTGCTAAAAAGAATGGTGAGCTACGTCATT			
Consensus (8401)		-----	-----	-----	-----
Section 177					
	(8449)	8449	8460	8470	8480
ClareAJ251507 (6600)		-----	-----	-----	-----
huNall18 (AK) (6090)		-----	-----	-----	-----
JeongAF225987 (8449)		ATTGAGCCAAAAGAATAAATTTTCATTTTTTATTGCATTTCACTTATTG			
Consensus (8449)		-----	-----	-----	-----
Section 178					
	(8497)	8497	8510	8520	8530
ClareAJ251507 (6600)		-----	-----	-----	-----
huNall18 (AK) (6090)		-----	-----	-----	-----
JeongAF225987 (8497)		GGCTCTGGGGTTTTTTGTTTTTGTGTTTTTGCTGTTGGCAGTTTAAAAT			
Consensus (8497)		-----	-----	-----	-----
Section 179					
	(8545)	8545	8550	8560	8570
ClareAJ251507 (6600)		-----	-----	-----	-----
huNall18 (AK) (6090)		-----	-----	-----	-----
JeongAF225987 (8545)		ATATATAATTAATAAAACCTGTGCTTGATCTGACATTTGTATACATAA			
Consensus (8545)		-----	-----	-----	-----
Section 180					
	(8593)	8593	8600	8610	8620
ClareAJ251507 (6600)		-----	-----	-----	-----
huNall18 (AK) (6090)		-----	-----	-----	-----
JeongAF225987 (8593)		AAGTTTACATGAATTTTACAACAACTAGTGCATGATTCACCAAGCAG			
Consensus (8593)		-----	-----	-----	-----
Section 181					
	(8641)	8641	8650	8660	8670
ClareAJ251507 (6600)		-----	-----	-----	-----
huNall18 (AK) (6090)		-----	-----	-----	-----
JeongAF225987 (8641)		TACTACAGAACAAAGGCAAATTAAGCAGCTTTGTGAACTTTTATGT			
Consensus (8641)		-----	-----	-----	-----
Section 182					
	(8689)	8689	8700	8710	8720
ClareAJ251507 (6600)		-----	-----	-----	-----
huNall18 (AK) (6090)		-----	-----	-----	-----
JeongAF225987 (8689)		GTGCAAAGGATCAAGTTCACATGTTCCAACCTTCAGGTTTGATAATAA			
Consensus (8689)		-----	-----	-----	-----

							Section 183
	(8737)	8737		8750	8760	8770	8784
ClareAJ251507	(6600)	-----					
huNall18 (AK)	(6090)	-----					
JeongAF225987	(8737)	TAGTAGTAACCACCTACAATAGCTTTCAATTTCAATTAACCTCCCTTGG					
Consensus	(8737)						
							Section 184
	(8785)	8785	8790	8800	8810	8820	8832
ClareAJ251507	(6600)	-----					
huNall18 (AK)	(6090)	-----					
JeongAF225987	(8785)	CTATAAGCATCTAAACTCATCTTCTTTCAATATAATTGATGCTATCTC					
Consensus	(8785)						
							Section 185
	(8833)	8833	8840	8850	8860	8870	8880
ClareAJ251507	(6600)	-----					
huNall18 (AK)	(6090)	-----					
JeongAF225987	(8833)	CTAATTACTTGGTGGCTAATAAATGTTACATTCTTTGTTACTTAAATG					
Consensus	(8833)						
							Section 186
	(8881)	8881	8890	8900	8910		8928
ClareAJ251507	(6600)	-----					
huNall18 (AK)	(6090)	-----					
JeongAF225987	(8881)	CATTATATAAACTCCTATGTATACATAAGGTATTAATGATATAGTTAT					
Consensus	(8881)						
							Section 187
	(8929)	8929	8940	8950	8960		8976
ClareAJ251507	(6600)	-----					
huNall18 (AK)	(6090)	-----					
JeongAF225987	(8929)	TGAGAAATTTATATTAACCTTTTTTTTCAAGAACCCTTGGATTTATGTGA					
Consensus	(8929)						
							Section 188
	(8977)	8977		8990	9000	9010	9024
ClareAJ251507	(6600)	-----					
huNall18 (AK)	(6090)	-----					
JeongAF225987	(8977)	GGTCAAAACCAAACCTCTTATTCTCAGTGGAAAACTCCAGTTGTAATGC					
Consensus	(8977)						
							Section 189
	(9025)	9025	9030	9040	9050	9060	9072
ClareAJ251507	(6600)	-----					
huNall18 (AK)	(6090)	-----					
JeongAF225987	(9025)	ATATTTTTTAAAGACAATTTGGATCTAAATATGTATTTTCATAATTCTCC					
Consensus	(9025)						

Section 190

	(9073)	9073	9080	9090	9100	9110	9120
ClareAJ251507 (6600)		-----	-----	-----	-----	-----	-----
huNall18 (AK) (6090)		-----	-----	-----	-----	-----	-----
JeongAF225987 (9073)		CATAATAAATTATATAAGGTGGAAAAAAAAAAAAAAAAAAAAAAAAAAAA					
Consensus (9073)							

Section 191

	(9121)	9123
ClareAJ251507 (6600)		---
huNall18 (AK) (6090)		---
JeongAF225987 (9121)		AAA
Consensus (9121)		

		Section 1				
		(1)	1	10	20	30
ClareAJ251507protein		(1)	MAQALLVPPGPESFRLFTRESLAAIEKRAAEEKAKKPKKE			
Translation of huNall18 (AK)		(1)	MAQALLVPPGPESFRLFTRESLAAIEKRAAEEKAKKPKKE			
Translation of JeongAF225987		(1)	MAQALLVPPGPESFRLFTRESLAAIEKRAAEEKAKKPKKE			
Consensus		(1)	MAQALLVPPGPESFRLFTRESLAAIEKRAAEEKAKKPKKE			
		Section 2				
		(41)	41	50	60	70
ClareAJ251507protein		(41)	QDNDDENKPKPNSDLEAGKNLPFIYGDIPPEMVSEPLEDL			
Translation of huNall18 (AK)		(41)	QDNDDENKPKPNSDLEAGKNLPFIYGDIPPEMVSEPLEDL			
Translation of JeongAF225987		(41)	QDNDDENKPKPNSDLEAGKNLPFIYGDIPPEMVSEPLEDL			
Consensus		(41)	QDNDDENKPKPNSDLEAGKNLPFIYGDIPPEMVSEPLEDL			
		Section 3				
		(81)	81	90	100	110
ClareAJ251507protein		(81)	DPYYINKKTFIVMNKGKAI FRFSATSALYILTPLNPVRKI			
Translation of huNall18 (AK)		(81)	DPYYINKKTFIVMNKGKAI FRFSATSALYILTPLNPVRKI			
Translation of JeongAF225987		(81)	DPYYINKKTFIVMNKGKAI FRFSATSALYILTPLNPVRKI			
Consensus		(81)	DPYYINKKTFIVMNKGKAI FRFSATSALYILTPLNPVRKI			
		Section 4				
		(121)	121	130	140	150
ClareAJ251507protein		(121)	AIKILVHSLFSMLIMCTILTNCVFMTLSNPPDWTKNVEYT			
Translation of huNall18 (AK)		(121)	AIKILVHSLFSMLIMCTILTNCVFMTLSNPPDWTKNVEYT			
Translation of JeongAF225987		(121)	AIKILVHSLFSMLIMCTILTNCVFMTLSNPPDWTKNVEYT			
Consensus		(121)	AIKILVHSLFSMLIMCTILTNCVFMTLSNPPDWTKNVEYT			
		Section 5				
		(161)	161	170	180	190
ClareAJ251507protein		(161)	FTGIYTFESLIKILARGFCLEDFTFLRDPWNWLD FSVIVM			
Translation of huNall18 (AK)		(161)	FTGIYTFESLIKILARGFCLEDFTFLRDPWNWLD FSVIVM			
Translation of JeongAF225987		(161)	FTGIYTFESLIKILARGFCLEDFTFLRDPWNWLD FSVIVM			
Consensus		(161)	FTGIYTFESLIKILARGFCLEDFTFLRDPWNWLD FSVIVM			
		Section 6				
		(201)	201	210	220	230
ClareAJ251507protein		(201)	AYVTEFVSLGNVSALRTFRVLRAKKTISVIPGLKTIVGAL			
Translation of huNall18 (AK)		(201)	AYVTEFVSLGNVSALRTFRVLRAKKTISVIPGLKTIVGAL			
Translation of JeongAF225987		(201)	AYVTEFVSLGNVSALRTFRVLRAKKTISVIPGLKTIVGAL			
Consensus		(201)	AYVTEFVSLGNVSALRTFRVLRAKKTISVIPGLKTIVGAL			
		Section 7				
		(241)	241	250	260	270
ClareAJ251507protein		(241)	IQSVKKLS DVMILTVFCLSVFALIGLQLFMGNLRNKCLQW			
Translation of huNall18 (AK)		(241)	IQSVKKLS DVMILTVFCLSVFALIGLQLFMGNLRNKCLQW			
Translation of JeongAF225987		(241)	IQSVKKLS DVMILTVFCLSVFALIGLQLFMGNLRNKCLQW			
Consensus		(241)	IQSVKKLS DVMILTVFCLSVFALIGLQLFMGNLRNKCLQW			

Section 8					
	(281)	281	290	300	310 320
ClareAJ251507protein	(281)	PPSDSAFETNTTSYFNGTMDSNGTFVNVTMSTFNWKDYIG			
Translation of huNall18 (AK)	(281)	PPSDSAFETNTTSYFNGTMDSNGTFVNVTMSTFNWKDYIG			
Translation of JeongAF225987	(281)	PPSDSAFETNTTSYFNGTMDSNGTFVNVTMSTFNWKDYIG			
Consensus	(281)	PPSDSAFETNTTSYFNGTMDSNGTFVNVTMSTFNWKDYIG			
Section 9					
	(321)	321	330	340	350 360
ClareAJ251507protein	(321)	DDSHFYVLDGQKDP LLCGNGSDAGQCPEGYICVKAGRNP			
Translation of huNall18 (AK)	(321)	DDSHFYVLDGQKDP LLCGNGSDAGQCPEGYICVKAGRNP			
Translation of JeongAF225987	(321)	DDSHFYVLDGQKDP LLCGNGSDAGQCPEGYICVKAGRNP			
Consensus	(321)	DDSHFYVLDGQKDP LLCGNGSDAGQCPEGYICVKAGRNP			
Section 10					
	(361)	361	370	380	390 400
ClareAJ251507protein	(361)	YGYTSFDTF SWAFLSLFRLMTQDYWENLYQLTLRAAGKTY			
Translation of huNall18 (AK)	(361)	YGYTSFDTF SWAFLSLFRLMTQDYWENLYQLTLRAAGKTY			
Translation of JeongAF225987	(361)	YGYTSFDTF SWAFLSLFRLMTQDYWENLYQLTLRAAGKTY			
Consensus	(361)	YGYTSFDTF SWAFLSLFRLMTQDYWENLYQLTLRAAGKTY			
Section 11					
	(401)	401	410	420	430 440
ClareAJ251507protein	(401)	MIFFVLVIFLGSFYLVNLILAVVAMAYEEQNQATLEEAEQ			
Translation of huNall18 (AK)	(401)	MIFFVLVIFLGSFYLVNLILAVVAMAYEEQNQATLEEAEQ			
Translation of JeongAF225987	(401)	MIFFVLVIFLGSFYLVNLILAVVAMAYEEQNQATLEEAEQ			
Consensus	(401)	MIFFVLVIFLGSFYLVNLILAVVAMAYEEQNQATLEEAEQ			
Section 12					
	(441)	441	450	460	470 480
ClareAJ251507protein	(441)	KEAEFQQMLEQLKKQQEEAQAVAAASAASRDFSGIGGLGE			
Translation of huNall18 (AK)	(441)	KEAEFQQMLEQLKKQQEEAQAVAAASAASRDFSGIGGLGE			
Translation of JeongAF225987	(441)	KEAEFQQMLEQLKKQQEEAQAVAAASAASRDFSGIGGLGE			
Consensus	(441)	KEAEFQQMLEQLKKQQEEAQAVAAASAASRDFSGIGGLGE			
Section 13					
	(481)	481	490	500	510 520
ClareAJ251507protein	(481)	LLESSSEASKLSSKSAKEWRNRRKKRRQREHLEGNNKGER			
Translation of huNall18 (AK)	(481)	LLESSSEASKLSSKSAKEWRNRRKKRRRREHLEGNNKGER			
Translation of JeongAF225987	(481)	LLESSSEASKLSSKGAKEWRNRRKKRRQREHLEGNNKGER			
Consensus	(481)	LLESSSEASKLSSKSAKEWRNRRKKRRQREHLEGNNKGER			
Section 14					
	(521)	521	530	540	550 560
ClareAJ251507protein	(521)	DSFPKSESEDSVKRSSFLFSMDGNRLTSDKKFCSPHQSL			
Translation of huNall18 (AK)	(521)	DSFPKSESEDSVKRSSFLFSMDGNRLTSDKKFCSPHQSL			
Translation of JeongAF225987	(521)	DSFPKSESEDSVKRSSFLFSMDGNRLTSDKKFCSPHQSL			
Consensus	(521)	DSFPKSESEDSVKRSSFLFSMDGNRLTSDKKFCSPHQSL			

Section 15

	(561)	561	570	580	590	600
ClareAJ251507protein	(561)	SIRGSLFSPRRNSKTSIFSFRGRAKDVGSENDFAADDEHST				
Translation of huNall18 (AK)	(561)	SIRGSLFSPRRNSKTSIFSFRGRAKDVGSENDFAADDEHST				
Translation of JeongAF225987	(561)	SIRGSLFSPRRNSKTSIFSFRGRAKDVGSENDFAADDEHST				
Consensus	(561)	SIRGSLFSPRRNSKTSIFSFRGRAKDVGSENDFAADDEHST				

Section 16

	(601)	601	610	620	630	640
ClareAJ251507protein	(601)	FEDSESRRDSLFPVPHRHGERRNS-----				
Translation of huNall18 (AK)	(601)	FEDSESRRDSLFPVPHRHGERRNSNVSQASMSSRMVPGLPA				
Translation of JeongAF225987	(601)	FEDGESRRDSLFPVPHRHGERRNSNVSQASMSSRMVPGLPA				
Consensus	(601)	FEDSESRRDSLFPVPHRHGERRNSNVSQASMSSRMVPGLPA				

Section 17

	(641)	641	650	660	670	680
ClareAJ251507protein	(624)	-----NGTTTETE				
Translation of huNall18 (AK)	(641)	NGKMHSTVDCNGVVSLVGGPSALTSTPTGQLPPEGTTTETE				
Translation of JeongAF225987	(641)	NGKMHSTVDCNGVVSLVGGPSALTSTPTGQLPPEGTTTETE				
Consensus	(641)	NGKMHSTVDCNGVVSLVGGPSALTSTPTGQLPPEGTTTETE				

Section 18

	(681)	681	690	700	710	720
ClareAJ251507protein	(632)	VRKRRLLSSYQISMEMLEDSSGRQRAVSIASILTNTMEELE				
Translation of huNall18 (AK)	(681)	VRKRRLLSSYQISMEMLEDSSGRQRAVSIASILTNTMEELE				
Translation of JeongAF225987	(681)	VRKRRLLSSYQISMEMLEDSSGRQRAVSIASILTNTMEELE				
Consensus	(681)	VRKRRLLSSYQISMEMLEDSSGRQRAVSIASILTNTMEELE				

Section 19

	(721)	721	730	740	750	760
ClareAJ251507protein	(672)	ESRQKCPCWCYRFANVFLIWDCCDAWLKVKHLVNLIVMDP				
Translation of huNall18 (AK)	(721)	ESRQKCPCWCYRFANVFLIWDCCDAWLKVKHLVNLIVMDP				
Translation of JeongAF225987	(721)	ESRQKCPCWCYRFANVFLIWDCCDAWLKVKHLVNLIVMDP				
Consensus	(721)	ESRQKCPCWCYRFANVFLIWDCCDAWLKVKHLVNLIVMDP				

Section 20

	(761)	761	770	780	790	800
ClareAJ251507protein	(712)	FVDLAITICIVLNTLFMAMEHYPMTEQFSSVLTVGNLVFT				
Translation of huNall18 (AK)	(761)	FVDLAITICIVLNTLFMAMEHYPMTEQFSSVLTVGNLVFT				
Translation of JeongAF225987	(761)	FVDLAITICIVLNTLFMAMEHYPMTEQFSSVLTVGNLVFT				
Consensus	(761)	FVDLAITICIVLNTLFMAMEHYPMTEQFSSVLTVGNLVFT				

Section 21

	(801)	801	810	820	830	840
ClareAJ251507protein	(752)	GIFTAEMVLKIIAMDPIYYYFQEGWNIFDGIIVSLSLMELG				
Translation of huNall18 (AK)	(801)	GIFTAEMVLKIIAMDPIYYYFQEGWNIFDGIIVSLSLMELG				
Translation of JeongAF225987	(801)	GIFTAEMVLKIIAMDPIYYYFQEGWNIFDGIIVSLSLMELG				
Consensus	(801)	GIFTAEMVLKIIAMDPIYYYFQEGWNIFDGIIVSLSLMELG				

Section 22					
	(841)	841	850	860	870 880
ClareAJ251507protein	(792)	LSNVEGLSVLRSFRLLRVFKLAKSWPTLNMLIKIIGNSVG			
Translation of huNall18 (AK)	(841)	LSNVEGLSVLRSFRLLRVFKLAKSWPTLNMLIKIIGNSVG			
Translation of JeongAF225987	(841)	LSNVEGLSVLRSFRLLRVFKLAKSWPTLNMLIKIIGNSVG			
Consensus	(841)	LSNVEGLSVLRSFRLLRVFKLAKSWPTLNMLIKIIGNSVG			
Section 23					
	(881)	881	890	900	910 920
ClareAJ251507protein	(832)	ALGNLTlVLAIIVFIFAVVGMQLFGKSYKECVCKINDDCT			
Translation of huNall18 (AK)	(881)	ALGNLTlVLAIIVFIFAVVGMQLFGKSYKECVCKINDDCT			
Translation of JeongAF225987	(881)	ALGNLTlVLAIIVFIFAVVGMQLFGKSYKECVCKINDDCT			
Consensus	(881)	ALGNLTlVLAIIVFIFAVVGMQLFGKSYKECVCKINDDCT			
Section 24					
	(921)	921	930	940	950 960
ClareAJ251507protein	(872)	LPRWHMNDFFHSFLIVFRVLCGEWIETMWDCMEVAGQTM			
Translation of huNall18 (AK)	(921)	LPRWHMNDFFHSFLIVFRVLCGEWIETMWDCMEVAGQTM			
Translation of JeongAF225987	(921)	LPRWHMNDFFHSFLIVFRVLCGEWIETMWDCMEVAGQTM			
Consensus	(921)	LPRWHMNDFFHSFLIVFRVLCGEWIETMWDCMEVAGQTM			
Section 25					
	(961)	961	970	980	990 1000
ClareAJ251507protein	(912)	LIVFMLVMVIGNLVVLNLFLLALLSSFSSDNLAATDDDDNE			
Translation of huNall18 (AK)	(961)	LIVFMLVMVIGNLVVLNLFLLALLSSFSSDNLAATDDDDNE			
Translation of JeongAF225987	(961)	LIVFMLVMVIGNLVVLNLFLLALLSSFSSDNLAATDDDDNE			
Consensus	(961)	LIVFMLVMVIGNLVVLNLFLLALLSSFSSDNLAATDDDDNE			
Section 26					
	(1001)	1001	1010	1020	1030 1040
ClareAJ251507protein	(952)	MNNLQIAVGRMQKGIDYVKNKMRECFQKAFFRKPKVIEIH			
Translation of huNall18 (AK)	(1001)	MNNLQIAVGRMQKGIDYVKNKMRECFQKAFFRKPKVIEIH			
Translation of JeongAF225987	(1001)	MNNLQIAVGRMQKGIDYVKNKMRECFQKAFFRKPKVIEIH			
Consensus	(1001)	MNNLQIAVGRMQKGIDYVKNKMRECFQKAFFRKPKVIEIH			
Section 27					
	(1041)	1041	1050	1060	1070 1080
ClareAJ251507protein	(992)	EGNKIDSCMSNNTGIEISKELNYLRDGNNGTTSGVGTGSSV			
Translation of huNall18 (AK)	(1041)	EGNKIDSCMSNNTGIEISKELNYLRDGNNGTTSGVGTGSSV			
Translation of JeongAF225987	(1041)	EGNKIDSCMSNNTGIEISKELNYLRDGNNGTTSGVGTGSSV			
Consensus	(1041)	EGNKIDSCMSNNTGIEISKELNYLRDGNNGTTSGVGTGSSV			
Section 28					
	(1081)	1081	1090	1100	1110 1120
ClareAJ251507protein	(1032)	EKYVIDENDYMSFINNPSLTVTVPPIAVGESDFENLNTEEF			
Translation of huNall18 (AK)	(1081)	EKYVIDENDYMSFINNPSLTVTVPPIAVGESDFENLNTEEF			
Translation of JeongAF225987	(1081)	EKYVIDENDYMSFINNPSLTVTVPPIAVGESDFENLNTEEF			
Consensus	(1081)	EKYVIDENDYMSFINNPSLTVTVPPIAVGESDFENLNTEEF			

Section 29

	(1121)	1121	1130	1140	1150	1160
ClareAJ251507protein (1072)		SSESELEESKEKLNATSSSEGSTVDVVLPREGEQAETEPE				
Translation of huNall18 (AK) (1121)		SSESELEESKEKLNATSSSEGSTVDVVLPREGEQAETEPE				
Translation of JeongAF225987 (1121)		SSESELEESKEKLNATSSSEGSTVDVVLPREGEQAETEPE				
Consensus (1121)		SSESELEESKEKLNATSSSEGSTVDVVLPREGEQAETEPE				

Section 30

	(1161)	1161	1170	1180	1190	1200
ClareAJ251507protein (1112)		EDLKPEACFTEGCIKKFPFCQVSTEEGKGKIWWNLRKTCY				
Translation of huNall18 (AK) (1161)		EDLKPEACFTEGCIKKFPFCQVSTEEGKGKIWWNLRKTCY				
Translation of JeongAF225987 (1161)		EDFKPEACFTEGCIKKFPFCQVSTEEGKGKIWWNLRKTCY				
Consensus (1161)		EDLKPEACFTEGCIKKFPFCQVSTEEGKGKIWWNLRKTCY				

Section 31

	(1201)	1201	1210	1220	1230	1240
ClareAJ251507protein (1152)		SIVEHNWFETFIVFMILLSSGALAFEDIYIEQRKTIKTM				
Translation of huNall18 (AK) (1201)		SIVEHNWFETFIVFMILLSSGALAFEDIYIEQRKTIKTM				
Translation of JeongAF225987 (1201)		SIVEHNWFETFIVFMILLSSGALAFEDIYIEQRKTIKTM				
Consensus (1201)		SIVEHNWFETFIVFMILLSSGALAFEDIYIEQRKTIKTM				

Section 32

	(1241)	1241	1250	1260	1270	1280
ClareAJ251507protein (1192)		EYADKVFTYIIFILEMLLKWVAYGFQTYFTNAWCWLDLFLIV				
Translation of huNall18 (AK) (1241)		EYADKVFTYIIFILEMLLKWVAYGFQTYFTNAWCWLDLFLIV				
Translation of JeongAF225987 (1241)		EYADKVFTYIIFILEMLLKWVAYGFQTYFTNAWCWLDLFLIV				
Consensus (1241)		EYADKVFTYIIFILEMLLKWVAYGFQTYFTNAWCWLDLFLIV				

Section 33

	(1281)	1281	1290	1300	1310	1320
ClareAJ251507protein (1232)		DVSLVSLVANALGYSELGAIKSLRTLRLRPLRLSRFEG				
Translation of huNall18 (AK) (1281)		DVSLVSLVANALGYSELGAIKSLRTLRLRPLRLSRFEG				
Translation of JeongAF225987 (1281)		DVSLVSLVANALGYSELGAIKSLRTLRLRPLRLSRFEG				
Consensus (1281)		DVSLVSLVANALGYSELGAIKSLRTLRLRPLRLSRFEG				

Section 34

	(1321)	1321	1330	1340	1350	1360
ClareAJ251507protein (1272)		MRVVVNALVGAIIPSIMNVLLVCLIFWLIFSIMGVNLFAGK				
Translation of huNall18 (AK) (1321)		MRVVVNALVGAIIPSIMNVLLVCLIFWLIFSIMGVNLFAGK				
Translation of JeongAF225987 (1321)		MRVVVNALVGAIIPSIMNVLLVCLIFWLIFSIMGVNLFAGK				
Consensus (1321)		MRVVVNALVGAIIPSIMNVLLVCLIFWLIFSIMGVNLFAGK				

Section 35

	(1361)	1361	1370	1380	1390	1400
ClareAJ251507protein (1312)		FYHCVNMTTGNMFDISDVNNLSDCQALGKQARWKNVKVNF				
Translation of huNall18 (AK) (1361)		FYHCVNMTTGNMFDISDVNNLSDCQALGKQARWKNVKVNF				
Translation of JeongAF225987 (1361)		FYHCVNMTTGNMFDISDVNNLSDCQALGKQARWKNVKVNF				
Consensus (1361)		FYHCVNMTTGNMFDISDVNNLSDCQALGKQARWKNVKVNF				

Section 36

	(1401)	1401	1410	1420	1430	1440
ClareAJ251507protein (1352)		DNVGAGYLALLQVATFKGWMDIMYAAVDSRDV	KLQPVYEE			
Translation of huNall18 (AK) (1401)		DNVGAGYLALLQVATFKGWMDIMYAAVDSRDV	KLQPVYEE			
Translation of JeongAF225987 (1401)		DNVGAGYLALLQVATFKGWMDIMYAAVDSRDV	KLQPVYEE			
Consensus (1401)		DNVGAGYLALLQVATFKGWMDIMYAAVDSRDV	KLQPVYEE			

Section 37

	(1441)	1441	1450	1460	1470	1480
ClareAJ251507protein (1392)		NLYMYLYFVIFIIFGSFFTLNLF	FIGVIIDN	FNQQKKK	FGG	
Translation of huNall18 (AK) (1441)		NLYMYLYFVIFIIFGSFFTLNLF	FIGVIIDN	FNQQKKK	FGG	
Translation of JeongAF225987 (1441)		NLYMYLYFVIFIIFGSFFTLNLF	FIGVIIDN	FNQQKKK	FGG	
Consensus (1441)		NLYMYLYFVIFIIFGSFFTLNLF	FIGVIIDN	FNQQKKK	FGG	

Section 38

	(1481)	1481	1490	1500	1510	1520
ClareAJ251507protein (1432)		QDIFMTEEQKKYYNAMKKLGSKKPQKPI	PRPANKFQGMVF			
Translation of huNall18 (AK) (1481)		QDIFMTEEQKKYYNAMKKLGSKKPQKPI	PRPANKFQGMVF			
Translation of JeongAF225987 (1481)		QDIFMTEEQKKYYNAMKKLGSKKPQKPI	PRPANKFQGMVF			
Consensus (1481)		QDIFMTEEQKKYYNAMKKLGSKKPQKPI	PRPANKFQGMVF			

Section 39

	(1521)	1521	1530	1540	1550	1560
ClareAJ251507protein (1472)		DFVTRQVFDISIMILICLNMVTMMVETDDQ	GKYMTLVLSR			
Translation of huNall18 (AK) (1521)		DFVTRQVFDISIMILICLNMVTMMVETDDQ	GKYMTLVLSR			
Translation of JeongAF225987 (1521)		DFVTRQVFDISIMILICLNMVTMMVETDDQ	GKYMTLVLSR			
Consensus (1521)		DFVTRQVFDISIMILICLNMVTMMVETDDQ	GKYMTLVLSR			

Section 40

	(1561)	1561	1570	1580	1590	1600
ClareAJ251507protein (1512)		INLVFIVLFTGEFVL	KLVS	LRHYYFTIGWNIFDFV	VVILS	
Translation of huNall18 (AK) (1561)		INLVFIVLFTGEFVL	KLVS	LRHYYFTIGWNIFDFV	VVILS	
Translation of JeongAF225987 (1561)		INLVFIVLFTGEFVL	KLVS	LRHYYFTIGWNIFDFV	VVILS	
Consensus (1561)		INLVFIVLFTGEFVL	KLVS	LRHYYFTIGWNIFDFV	VVILS	

Section 41

	(1601)	1601	1610	1620	1630	1640
ClareAJ251507protein (1552)		IVGMFLAEMIEKYFVSPTLFRVIRLARIGRIL	RRIKGAKG			
Translation of huNall18 (AK) (1601)		IVGMFLAEMIEKYFVSPTLFRVIRLARIGRIL	RRIKGAKG			
Translation of JeongAF225987 (1601)		IVGMFLAEMIEKYSVSPTLFRVIRLARIGRIL	RRIKGAKG			
Consensus (1601)		IVGMFLAEMIEKYFVSPTLFRVIRLARIGRIL	RRIKGAKG			

Section 42

	(1641)	1641	1650	1660	1670	1680
ClareAJ251507protein (1592)		IRTLFLALMMSLPALFNIGLLLFLVMFIYAIFG	MSNFAYV			
Translation of huNall18 (AK) (1641)		IRTLFLALMMSLPALFNIGLLLFLVMFIYAIFG	MSNFAYV			
Translation of JeongAF225987 (1641)		IRTLFLALMMSLPALFNIGLLLFLVMFIYAIFG	MSNFAYV			
Consensus (1641)		IRTLFLALMMSLPALFNIGLLLFLVMFIYAIFG	MSNFAYV			

Section 43

	(1681)	1681	1690	1700	1710	1720
ClareAJ251507protein (1632)		KKEAGIDDMFNFETFGNSMICLFQITTSAGWDGLLAPILN				
Translation of huNall18 (AK) (1681)		KKEAGIDDMFNFETFGNSMICLFQITTSAGWDGLLAPILN				
Translation of JeongAF225987 (1681)		KKEAGIDDMFNFETFGNSMICLFQITTSAGWDGLLAPILN				
Consensus (1681)		KKEAGIDDMFNFETFGNSMICLFQITTSAGWDGLLAPILN				

Section 44

	(1721)	1721	1730	1740	1750	1760
ClareAJ251507protein (1672)		SAPPDCDPDTIHPGSSVKGDCG N PSVGIFFFVSYIIISFL				
Translation of huNall18 (AK) (1721)		SAPPDCDPDTIHPGSSVKGDCG N PSVGIFFFVSYIIISFL				
Translation of JeongAF225987 (1721)		SAPPDCDPDTIHPGSSVKGDRGDPVGIFFFVSYIIISFL				
Consensus (1721)		SAPPDCDPDTIHPGSSVKGDCG N PSVGIFFFVSYIIISFL				

Section 45

	(1761)	1761	1770	1780	1790	1800
ClareAJ251507protein (1712)		VVVNMYIAVILENFSVATEESAEP L SEDDFEMFYEVWEKF				
Translation of huNall18 (AK) (1761)		VVVNMYIAVILENFSVATEESAEP L SEDDFEMFYEVWEKF				
Translation of JeongAF225987 (1761)		VVVNMYIAVILENFSVATEESAEP L SEDDFEMFYEVWEKF				
Consensus (1761)		VVVNMYIAVILENFSVATEESAEP L SEDDFEMFYEVWEKF				

Section 46

	(1801)	1801	1810	1820	1830	1840
ClareAJ251507protein (1752)		DPDATQFIEFSKLSDFAAALDPPL L IAKPNKVQLIAMDL P				
Translation of huNall18 (AK) (1801)		DPDATQFIEFSKLSDFAAALDPPL L IAKPNKVQLIAMDL P				
Translation of JeongAF225987 (1801)		DPDATQFIEFSKLSDFAAALDPPL L IAKPNKVQLIAMDL P				
Consensus (1801)		DPDATQFIEFSKLSDFAAALDPPL L IAKPNKVQLIAMDL P				

Section 47

	(1841)	1841	1850	1860	1870	1880
ClareAJ251507protein (1792)		MVSGDRIHCLDILFAFTKRV L GESGEMDALRIQMEDRFMA				
Translation of huNall18 (AK) (1841)		MVSGDRIHCLDILFAFTKRV L GESGEMDALRIQMEDRFMA				
Translation of JeongAF225987 (1841)		MVSGDRIHCLDILFAFTKRV L CESGEMDALRIQMEDRFMA				
Consensus (1841)		MVSGDRIHCLDILFAFTKRV L GESGEMDALRIQMEDRFMA				

Section 48

	(1881)	1881	1890	1900	1910	1920
ClareAJ251507protein (1832)		SNPSKVSYPEITTT L KRKQEEVSA A IIQRNFRCYLLKQRL				
Translation of huNall18 (AK) (1881)		SNPSKVSYPEITTT L KRKQEEVSA A IIQRNFRCYLLKQRL				
Translation of JeongAF225987 (1881)		SNPSKVSYPEITTT L KRKQEEVSA A IIQRNFRCYLLKQRL				
Consensus (1881)		SNPSKVSYPEITTT L KRKQEEVSA A IIQRNFRCYLLKQRL				

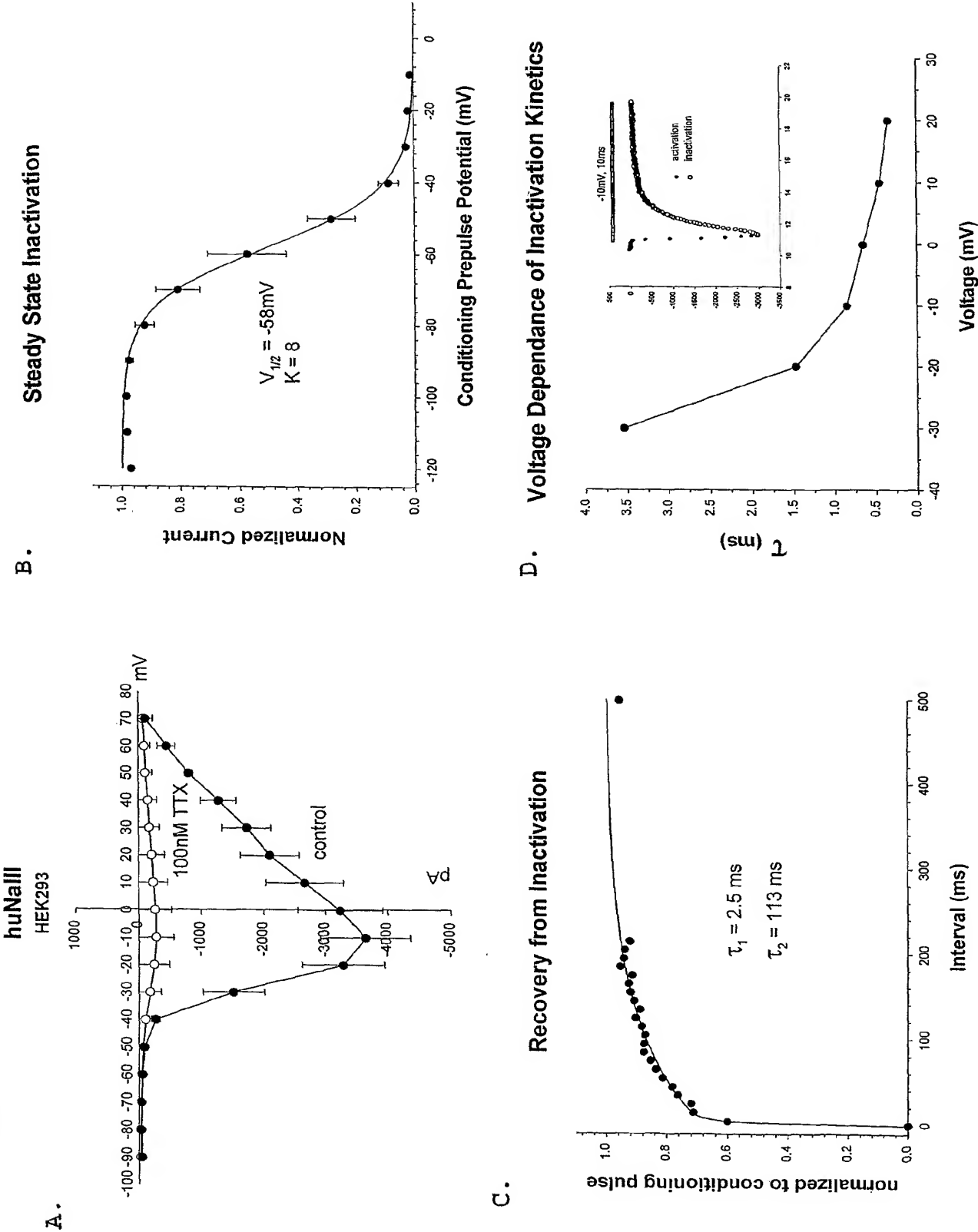
Section 49

	(1921)	1921	1930	1940	1950	1960
ClareAJ251507protein (1872)		KNISSN N YNKEAIKGRIDLP I KQDMIIDKLNGNSTPEKTDG				
Translation of huNall18 (AK) (1921)		KNISSN N YNKEAIKGRIDLP I KQDMIIDKLNGNSTPEKTDG				
Translation of JeongAF225987 (1921)		KNISSN N YNKEAIKGRIDLP I KQDMIIDKLNGNSTPEKTDG				
Consensus (1921)		KNISSN N YNKEAIKGRIDLP I KQDMIIDKLNGNSTPEKTDG				

Section 50

	(1961)	1961	1970	1980	1990	2000
ClareAJ251507protein (1912)	SSSTT	S	PPSYDSVTKPDKEKFEKDKPEKESKGKEVRENQK			
Translation of huNall118 (AK) (1961)	SSSTT	S	PPSYDSVTKPDKEKFEKDKPEKESKGKEVRENQK			
Translation of JeongAF225987 (1961)	SSSTT	PPSYDSVTKPDKEKFEKDKPEKESKGKEVRENQK				
Consensus (1961)	SSSTT	S	PPSYDSVTKPDKEKFEKDKPEKESKGKEVRENQK			

Figure 3



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GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR,
KZ, LC, LK, LR, LS, LT, LU, LV, MA, MD, MG, MK, MN,
MW, MX, MZ, NI, NO, NZ, OM, PG, PH, PL, PT, RO, RU,
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(54) Title: SPLICE VARIANT OF HUMAN SODIUM III CHANNEL (HNAIII18)

(57) Abstract: Described herein is a splice variant of the human NaIII channel α subunit, designated hNaIII18. Also described are nucleotide and amino acid sequence for hNaIII18, oligonucleotide primers and probes for hNaIII18, hNaIII18 regulatory sequences, hNaIII18-specific antibodies, methods of detecting hNaIII18 proteins or nucleic acids, and methods of screening for modulators of hNaIII18 expression or activity.



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INTERNATIONAL SEARCH REPORT

International application No.

PCT/US03/38796

A. CLASSIFICATION OF SUBJECT MATTER

IPC(7) : C12 Q 1/68

US CL : 435/6, 320.1, 325, 455, 91.41; 536/23.1, 23.2, 23.5, 24.3, 24.31

According to International Patent Classification (IPC) or to both national classification and IPC

B. FIELDS SEARCHED

Minimum documentation searched (classification system followed by classification symbols)

U.S. : 435/6, 320.1, 325, 455, 91.41; 536/23.1, 23.2, 23.5, 24.3, 24.31

Documentation searched other than minimum documentation to the extent that such documents are included in the fields searched

Electronic data base consulted during the international search (name of data base and, where practicable, search terms used)

Please See Continuation Sheet

C. DOCUMENTS CONSIDERED TO BE RELEVANT

Category *	Citation of document, with indication, where appropriate, of the relevant passages	Relevant to claim No.
Y	US 6,110,672 A (MANDEL et al.) 29 August 2000 (29.08.2000), especially Examples.	1-21

☐ Further documents are listed in the continuation of Box C.

☐ See patent family annex.

* Special categories of cited documents:

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INTERNATIONAL SEARCH REPORT

PCT/US03/38796

Continuation of B. FIELDS SEARCHED Item 3:
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search terms: nucleotide, polypeptide, sodium channel, human.